

DEPARTMENT OF THE NAVY

OFFICE OF THE GENERAL COUNSEL

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Navy Case No. 78,824

Commissioner of Patents and Trademarks Washington, D. C. 20231

Applicant(s): James H. Adams, Jr., Paul Boberg, Buddy Brownstein, William Dietrich, Erwin Flueckiger, Edward Petersen, Margaret Shea, Don Smart, and Edward Smith

For: METHOD AND APPARATUS FOR MODELING COSMIC RAY EFFECTS ON MICROELECTRONICS

Sir:

Transmitted herewith are the papers above-identified constituting a Patent Application filed by the Department of the Navy on behalf of the above-names Applicant(s).

The total filing fee has been computed in accordance with the following formula:

Basic Fee -----\$790.00

Additional Fees:

- 1. Total number of claims in excess of 20 times \$22.00-----\$____
- 2. Number of independent claims minus 3 times \$82.00-----\$246.00

Total Filing Fee-----\$ 1036.00

Kindly charge the aforementioned total filing fees and any additional fees due, or credit overpayment of fees, to Deposit Account No. 04-0814.

Respectfully,

THOMAS E. McDONNELL

Reg. #26950

Encl:



DEPARTMENT OF THE NAVY

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Navy Case No. 78,824

COUNSEL FOR THE NAVAL RESEARCH LABORATORY WASHINGTON, D.C. 20375-5325

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re application: John H. Adams, et al.

For: METHOD AND APPARATUS FOR MODELING COSMIC RAY EFFECTS ON MICROELECTRONICS

NOTICE OF FILING WITHOUT OATH OR DECLARATION BY APPLICANT

Honorable Commissioner of Patents and Trademarks Washington, D.C. 20231

Sir:

Please take notice that this application is being filed pursuant to 37 C.F.R. 153(d) without the oath or declaration of the Applicant. This is being done to secure an early filing date.

Upon the Notice from the Patent Office required by 1.53(d), Applicant will file the required oath or declaration, and pay the surcharge as set forth in 37 C.F.R. 1.16(e).

Respectfully submitted,

THOMAS E. McDONNELL

Req. No. 26950

Date:



DEPARTMENT OF THE NAVY

OFFICE OF THE GENERAL COUNSEL

Navy Case No. 78,824 COUNSEL FOR THE NAVAL RESEARCH LABORATORY WASHINGTON, D.C. 20375-5325

APPLICATION FOR LETTERS PATENT

TO ALL WHOM IT MAY CONCERN:

BE IT KNOWN THAT ______ James H. Adams, Jr., Paul Boberg, Buddy Browstein, William Dietrich, Erwin Flueckiger, Edward Petersen, Margaret Shea, Don Smart, Edward Smith and Allen Tylka are citizens of the United States of America, and residents of __Alexandria, Va, College Park, Md, Oxen Hill, Md, Chicago, IL, Bern Switzerland, Fairfax, Va, Bedford, Ma, Bedford, Ma, Mounteagle, TN, Silver Spring, Md, have invented certain new and useful improvements in METHOD AND APPARATUS FOR MODELING COSMIC RAY EFFECTS ON MICROELECTRONICS of which the following is a specification:

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METHOD AND APPARATUS FOR MODELING COSMIC RAY EFFECTS ON MICROELECTRONICS

BACKGROUND OF THE INVENTION

Field of the Invention

The present invention relates generally to a method and apparatus for modeling the effects of cosmic rays on microelectronics in earth orbit, and more particularly to an improved method and apparatus for modeling these effects, these improvements reflecting a simpler, easier way for the user to operate the model over the internet, and more accurate modeling of these effects.

Description of the Related Art

For electronic components onboard satellites in earth orbit, exposure to cosmic rays represents a serious risk, due to the capacity of cosmic rays to induce single event effects (SEE) in these components. *See generally* Sherra E. Kerns, *Transient-Ionization and Single-Event Phenomena, in* IONIZING RADIATION EFFECTS IN MOS DEVICES AND CIRCUITS 485-91 (John Wiley & Sons, Inc., T.P. Ma et al. eds., 1989). In brief, SEEs occur when an energetic particle changes a particular device in an integrated circuit, thereby causing an error. To date, the only effective methods for hardening these circuits against SEEs have been shielding, redundancy, and error detection and correction (EDAC).

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Unfortunately, these measures are not always effective, and the failure of a single component may lead to the total loss of a multi-million dollar satellite. With the potential loss associated with this risk so high it is desired to have a way to accurately predict the magnitude of this risk, as well as a way of predicting how successful ameliorative efforts are likely to be.

CREME (short for Cosmic Ray Effects on Microelectronics) was a software package developed by the Naval Research Laboratory in 1981 for modeling how a given electronic chip on a given satellite with a given orbit and amount of shielding would hold up against cosmic ray bombardment. *See* Cosmic Ray Effects on Microelectronics, Part I: The Near-Earth Particle Environment, Adams et al., NRL Memorandum Report 4506; Cosmic Ray Effects on Microelectronics Part II: The Geomagnetic Cutoff Effects, Adams et al., NRL Memorandum Report 5099; Cosmic Ray Effects on Microelectronics, Part IV, Adams, NRL Memorandum Report 5901, each incorporated herein by reference, in their entireties, for all purposes. CREME had several shortcomings. It has been discovered that many of the predictions of CREME were inaccurate. Particular shortcomings of the original CREME software included its inaccurate modeling of the transmission of cosmic rays through earth's magnetosphere, and inaccurate modeling of the flux of heavy ions associated with solar flares.

Moreover, the implementation of this program was less than optimal, having a difficult user interface, and requiring each user to install, maintain, and run the software.

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Inventor(s): Adams et al.

NOTE ON NOMENCLATURE USED HEREIN

Some portions of the following disclosure describe method elements such as determining, selecting, dividing, processing, computing, calculating, numerically integrating, applying a function, displaying, and the like, and structure elements such as tables, memories, internet connections, and the like. These descriptions are the means used by persons of ordinary skill in the art of data processing to most effectively convey the substance of their work to other skilled artisans. Such methods and structures are intended to describe methods and structures for carrying out a set of steps on at least one programmed digital computer to reach a desired result. Thus, each of these steps requires a physical manipulation of concrete quantities, generally in the form of electrical, optical, and magnetic signals capable of being stored, retrieved, combined, and otherwise manipulated in such a programmed digital computer.

Accordingly, unless indicated otherwise, skilled artisans will recognize that as used herein, terms such as determining, selecting, dividing, processing, computing, calculating, numerically integrating, applying a function, displaying, and the like, refer to the operations of a programmed digital computer system, or similar electronic computing device, that manipulates and transforms data represented as physical quantities within the computer.

SUMMARY OF THE INVENTION

Accordingly, it is an object of this invention to improve the modeling of the effects of cosmic rays on microelectronics.

It is a further object of this invention to improve the modeling of solar heavy ion flux.

It is a further object of this invention to improve the modeling of geomagnetic transmission.

It is a further object of this invention to improve the user interface, by implementing an internet-based interface, allowing users to access a world wide web site connected to a server where the software is installed and maintained.

These and additional objects of the invention are accomplished by the structures and processes hereinafter described.

An aspect of the present invention is a method and apparatus for computing a geomagnetic transmission function. This apparatus includes a programmed digital computer running modeling software for modeling the transmission of cosmic ray particles through the magnetosphere. The software includes a model representing a solution to the Lorentz equation in a magnetic field given by:

$$\boldsymbol{B} = \boldsymbol{B}_{\text{IGRF}}(\boldsymbol{r},\!t') + \boldsymbol{B}_{\text{TSYG}}(\boldsymbol{K}\boldsymbol{p},\!\boldsymbol{r},\!t')$$

where \mathbf{B} is the earth's magnetic field, where $\mathbf{B}_{IGRF}(\mathbf{r},t')$ is the International Geomagnetic Reference Field model of earth's magnetic field, a standard internationally recognized representation of earth's magnetic field, and where $\mathbf{B}_{TSYG}(Kp,\mathbf{r},t')$ is the model of earth's magnetic field published by Tsyganenko in 1989, as modified by the inventors. $\mathbf{B}_{IGRF}(\mathbf{r},t')$ and $\mathbf{B}_{TSYG}(Kp,\mathbf{r},t')$ are discussed in further detail *infra*.

Another aspect of the present invention is a method and apparatus for computing a flux of particles at the outer surface of a satellite comprising, *inter alia*, an improved method and apparatus for computing a flux of solar heavy ions. This apparatus includes a programmed digital computer

of a satellite.

Another aspect of the invention is a method and apparatus comprising a programmed digital computer running modeling software for modeling the effect of cosmic rays on microelectronics, where this software embodies at least one of the two foregoing aspects of the invention.

running modeling software for modeling the flux of cosmic ray particles through the outer surface

Another aspect of the invention is a preferred embodiment of a method and apparatus comprising a programmed digital computer running modeling software for modeling the effect of cosmic rays on microelectronics, where this software embodies at least one of the two foregoing aspects of the invention, where this preferred embodiment is connected to a network, typically the internet, to permit remote users to use the invention.

BRIEF DESCRIPTION OF THE DRAWINGS

A more complete appreciation of the invention will be obtained readily by reference to the following Description of the Preferred Embodiments and the accompanying drawings in which like numerals in different figures represent the same structures or elements, wherein:

- FIG. 1 is a partial flowchart for a preferred embodiment of the invention.
- FIG. 2 is an elevation of the earth, showing the path of a cosmic ray particle in the magnetosphere.
- FIG. 3 is a flowchart for a method for calculating a geomagnetic transmission function for cosmic rays according to the invention.

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- FIG. 4 is a plot of a geomagnetic transmission function for a low inclination space shuttle orbit under quiet magnetospheric conditions.
- FIG. 5 is a plot of a geomagnetic transmission function for a lo inclination space shuttle orbit under stormy magnetospheric conditions.
- FIG. 6 is a plot of a geomagnetic transmission function for a space station orbit under quiet magnetospheric conditions.
- FIG. 7 is a plot of a geomagnetic transmission function for a space station orbit under stormy magnetospheric conditions.
- FIGS. 8 and 9 together are a flowchart for a method for modeling solar heavy ion flux according to the present invention.
- FIG. 10 shows a main menu for a user to operate the preferred CREME96 software through the internet.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

The following are each incorporated herein by reference, in their entireties, for all purposes:

- (A) "CREME96: A Revision of the Cosmic Ray Effects on Micro-Electronics Code", Transactions on Nuclear Science 44(6) 2150-60, Tylka et al. (1997);
- (B) "A Magnetosphereic Magnetic Field Model with a Warped Tail Current Sheet", Planet. Space Sci. 37(1) 5-20, Tsyganenko (1989).

Overview

In a preferred embodiment, the invention comprises several computer software modules running on a powerful server computer. Although this computer may be operated in a standalone mode, it is preferred to have this server connected to a network (typically the internet), so that remote users can operate the software, and store their resulting data either locally on the server or remotely on their own or another machine.

Overall, the software will include one or both of a geomagnetic transmission modeling module and a particle flux modeling module. Optionally, the software includes both of these modules, and also optionally the software includes one or more of several optional modules, including a shielding transport modeling module, a linear energy transfer modeling module, a protoninduced single event upset (SEU, a particular type of SEE, but unless noted otherwise used interchangeably herein with SEE) rate modeling module, and a direct ionization induced single event upset rate modeling module.

Examples of each of these modules have been produced, and will be discussed in greater detail below. The exemplary module for modeling geomagnetic transmission is referred to herein as GTRN. The exemplary module for modeling particle flux is referred to herein as FLUX. The exemplary module for modeling transport of particles through solid shielding is referred to herein as TRANS. The exemplary module for modeling linear energy transfer to a target structure (typically an integrated circuit) is referred to herein as LETSPEC. The exemplary module for modeling protoninduced SEU event upset rates in specified devices is referred to herein as PUP. The exemplary module for modeling SEU rates induced by direct ionization is referred to herein as HUP.

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Exemplary FORTRAN code for these exemplary modules appears infra. Together, the software for these exemplary modules is referred to herein as CREME96.

Referring to FIG. 1, one sees that CREME96 produces two types of outputs: proton-induced SEU rates, and direct ionization induced SEU rates. Each of these will represent an upset rate for a given electronic device in a satellite with a given shielding, under given environmental conditions, for a satellite in a given orbit or orbit segment.

Referring to 100, in the case where the specified satellite orbit is at or above a given altitude, nominally geosynchronous, the software will evaluate the fluxes of SEU-inducing particles at the outer surface of the satellite, from a number of sources including galactic cosmic rays (GCR), anomalous cosmic rays (ACR), and solar heavy ions. The environment at such satellite orbits is referred to herein as the "non-trapped energetic particle environment", since it is essentially equivalent to the environment at other locations that are at about 1 AU from the sun, but outside of the volume where trapped or quasi-trapped particles would be found in significant numbers.

Referring to 200, in the case where the specified satellite orbit is below a given altitude, nominally geosynchronous, before the flux at the outer surface of the satellite is evaluated, the shielding effect of the earth's magnetosphere is evaluated. As persons of ordinary skill in the art will recognize, the earth's magnetic field will shield out a significant fraction of charged particles. Thus, it is necessary to first subtract from the flux in the non-trapped energetic particle environment that fraction of the flux that will be shielded by the magnetosphere.

Referring to 300, the external hull of a satellite (typically made from thin aluminum panels) will shield the electronics of the satellite from a fraction of SEU-inducing particles. Thus, determining an SEU rate for microelectronics on a given satellite will require taking this shielding into account by correcting the flux for the energy lost in said shielding and for the fragmentation of the incident particles within said shielding, and subtracting from the flux of SEU-inducing particles the subset of SEU-inducing particles with energies that are not sufficient to penetrate through the hull of the satellite. This module of the software performs that function. Although, as indicated by the dashed lines in the flowchart, this module may be bypassed, it is highly preferred to *not* bypass this module.

Referring to 400, although an approximation of the satellite hull shielding may be obtained by assuming that the shielding is uniformly distributed about the electronics of interest, more realistic results may be obtained by specifying the particular distribution of shielding about the electronics for a particular satellite. This module of the software performs that function.

Referring to 500, after the software measures the transmission of SEU-inducing particles, the software will measure the effect of SEU-inducing protons on the electronics of interest.

Referring to 600, after the software measures the transmission of SEU-inducing particles, the software will calculate the linear energy transfer to the electronics of interest. This module of the software takes an input file in the form of a particle flux vs. kinetic energy table for each elemental species, and generates an output file in the form of a function specifying the combined flux of all

these species as a function of the rate of energy deposition. This is an intermediate result for the calculation of SEU rates attributable to direct ionization.

Referring to 700, this program module takes the output linear energy transfer spectrum from 600, combines this with parameters for the particular electronic device, calculates a SEU rate for the device attributable to direct ionization. The parameters are typically based on ground tests of the sensitivity to a particular chip to SEU-inducing radiation.

Having given an overview of this preferred embodiment of the invention, a more detailed description of the particular software modules making up this preferred embodiment is given.

Geomagnetic Transmission Function

Although it has been known that the magnetosphere shields cosmic rays, quantifying this effect to a degree of accuracy that could lead to accurate models of the effects of cosmic rays on electronics within the magnetosphere has proven elusive. One difficulty has been the lack of accurate models for the magnetosphere. Another difficulty has been that even if an accurate model of the magnetosphere were available, modeling the shielding effect of the magnetosphere requires an inordinate amount of computer processing time.

Referring to FIG. 2, this figure shows a representative path of a cosmic ray through the earth's magnetosphere (not shown). This path is highly irregular, including numerous changes of direction, and covering widely-spaced areas over earth's surface. Determining whether a particle

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with such a convoluted path will intersect a body as small as a typical satellite is a considerable challenge.

B_{IGRF} is the International Geomagnetic Reference Field promulgated by the International Association of Geomagnetism and Aeronomy. It is the internationally accepted standard expression for the earth's internal magnetic field. It has been discovered that the \mathbf{B}_{IGRF} is insufficient, and that the most well-known model of the magnetospheric B fields, the model provided by Tsyganenko et al., "A Magnetospheric Magnetic Field Model with a Warped Tail Current Sheet", Planet. Space Sci., 37(1) 5-20 (1989), is likewise insufficient. Corrections to this model are included in the present invention.

It has been discovered that within the earth's magnetosphere, the total magnetic field can be modeled by;

$$\mathbf{B} = \mathbf{B}_{IGRF}(\mathbf{r},t') + \mathbf{B}_{TSYG}(\mathbf{K}\mathbf{p},\mathbf{r},t')$$

where \mathbf{B}_{IGRF} is the International Geomagnetic Reference Field promulgated by the International Association of Geomagnetism and Aeronomy, and \mathbf{B}_{TSYG} is the modified Tsyganenko field given by $\mathbf{B}_{\mathrm{Xsm}}^{(M)} + \mathbf{B}_{\mathrm{Ysm}}^{(M)} + \mathbf{B}_{\mathrm{Zsm}}^{(M)}$, where coordinates in the solar magnetospheric system are denoted sm, and where coordinates in the solar magnetic coordinate system are denoted mg; and where

$$\mathbf{B}_{\mathsf{Xsm}}^{(\mathsf{T})} = \mathbf{Q}_{\mathsf{t}} \, \mathbf{X}_{\mathsf{sm}} \, \mathbf{Z}_{\mathsf{r}};$$

$$\mathbf{B}_{\mathsf{Ysm}}^{(\mathsf{T})} = \mathbf{Q}_{\mathsf{t}} \, \mathsf{y}_{\mathsf{sm}} \, \mathsf{z}_{\mathsf{r}};$$

 $\mathbf{B}_{z_{sm}}(T) = \frac{W(x,y)}{S_{T}} \left(C_{1} + C_{2} \frac{a_{T} + \xi_{T}}{S_{T}^{2}} \right) + \frac{x \frac{\partial W}{\partial x} + y \frac{\partial W}{\partial y}}{S_{T} + a_{T} + \xi_{T}} \times$

$$S_{T} = \begin{pmatrix} C_{T} + C_{T} & S_{T}^{2} \end{pmatrix}^{T} S_{T} + a_{T} + \xi_{T}$$

$$(C_{1} + \frac{C_{2}}{S_{T}}) + \mathbf{B}_{X}(T) \frac{\partial \mathbf{z}_{s}}{\partial \mathbf{x}} + \mathbf{B}_{Y}(T) \frac{\partial \mathbf{z}_{s}}{\partial \mathbf{y}} - Q_{T}D_{T} \left(\mathbf{x} \frac{\partial D_{T}}{\partial \mathbf{x}} + \mathbf{y} \frac{\partial D_{T}}{\partial \mathbf{y}} \right)$$

where W(x,y) = 0.5
$$\left[1 - \frac{x - x_0}{\left[\left(x - x_0\right)^2 + D_x^2\right]^{\frac{1}{2}}}\right]$$

where
$$Q_T = \frac{W(x,y)}{\xi_T S_T} \left[\frac{C_1}{S_T + a_T + x_T} + \frac{C_2}{S_T^2} \right];$$

where $z_r = z - z_s(x, y, \psi)$,

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$$z_{s}(x, y, \psi) = 0.5 \tan \psi \left(x + R_{c} - \sqrt{\left(x + R_{c} \right)^{2} + 16} \right)$$

$$- G \sin \psi \cdot y^{4} \left(y^{4} + L_{y}^{4} \right)^{-1},$$
where
$$S_{T,RC} = \sqrt{\rho^{2} + \left(a_{T,RC} + \xi_{T,RC} \right)^{2}},$$

$$\xi_{T,RC} = \sqrt{z_{r}^{2} + D_{T,RC}^{2}},$$

$$D_{T} = D_{0} + \delta y^{2} + \gamma_{T} h_{T}(x) + \gamma_{1} h_{1}(x),$$

when $Kp = 2^-, 2, 2^+$, $C_1 = -70.12$ when $Kp = 3^-, 3, 3^+$, $0, 0^+$, $C_1 = -162.5$ when $Kp = 4^-, 4, 4^+$, $C_1 = -128.4$ when $Kp \ge 5^-$, where $C_2 = -10014$ when $Kp = 0, 0^-$, $C_2 = -12800$ when $Kp = 1^-, 1, 1^+$, $C_2 = -14588$ when $Kp = 2^-, 2, 2^+$, $C_2 = -16125$ when $Kp = 3^-, 3, 3^+$, $0, 0^+$, $C_2 = -15806$ when $Kp = 4^-, 4, 4^+$, $C_2 = -16184$ when $Kp \ge 5^-$, where $a_T = 13.55$ when $Kp = 0, 0^+$, $a_T = 13.81$ when $Kp = 1^-, 1, 1^+$, $a_T = 15.08$ when $Kp = 2^-, 2, 2^+$, $a_T = 15.63$ when $Kp = 3^-, 3, 3^+$, $0, 0^+$, $a_T = 16.11$ when $Kp = 4^-, 4, 4^+$, $a_T = 15.85$ when $Kp \ge 5^-$,

where $C_1 = -98.72$ when $Kp = 0, 0^-, C_1 = -35.64$ when $Kp = 1^-, 1, 1^+, C_1 = -77.45$

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where D_0 = 2.08 when Kp = 0, 0^+ , D_0 = 1.664 when Kp = 1^- ,1,1 $^+$, D_0 = 1.541 when Kp = 2^- ,2,2 $^+$, D_0 = 0.9351 when Kp = 3^- ,3,3 $^+$, 0, 0^+ , D_0 = 0.7677 when Kp = 4^- ,4,4 $^+$, D_0 = 0.3325 when $Kp \ge 5^-$, where R_c = 8 R_E and L_v = 10 R_E ,

$$\mathbf{B}_{\mathbf{X}}^{(\mathbf{RC})} = \mathbf{Q}_{\mathbf{RC}} \mathbf{X} \mathbf{Z}_{\mathbf{r}};$$

$$\mathbf{B}_{\mathbf{Y}}^{(RC)} = \mathbf{Q}_{RC} \mathbf{y} \mathbf{z}_{r};$$

$$\mathbf{B}_{z}^{(RC)} = C_{5} \frac{2(\mathbf{a}_{RC} + \xi_{RC})^{2} - \rho^{2}}{S_{RC}^{5}} + B_{x}^{RC} \frac{\partial z_{s}}{\partial x} + B_{y}^{RC} \frac{\partial z_{s}}{\partial y} - Q_{RC}D_{RC}x \frac{\partial D_{RC}}{\partial x};$$

where
$$Q_{RC} = 3C_5 \xi_{RC}^{-1} S_{RC}^{-5} (a_{RC} + \xi_{RC})$$

 $D_{RC} = D_0 + \gamma_{RC} h_{RC}(x) + \gamma_1 h_1(x)$
 $h_{T,RC} = 0.5[1 + x(x^2 +)$
 $h_{T,RC} = 0.5[1 + x(x^2 + L_{T,RC}^2)^{-1/2}],$
 $h_1 = 0.5\{1 - (x + 16)[(x + 16)^2 + 36]^{-1/2}\},$

$$C_5(Dst) = -10220 + 408.5 \cdot Dst$$

$$\mathbf{B}_{XYZ}^{(C)} = C_3(F^+_{x,y,z} + F^-_{x,y,z}) + C_4(F^+_{x,y,z} - F^-_{x,y,z}), \text{ where}$$

$$\begin{split} \left\{ \frac{F_{x}^{\pm}}{F_{y}^{\pm}} \right\} &= \pm \frac{W_{c}(x,y)}{S^{\pm}[S^{\pm}\pm(z\pm R_{T})]} \times \left\{ \frac{x}{y} \right\}, \\ F_{z}^{\pm} &= \frac{W_{c}(x,y)}{S^{\pm}} + \left(x \frac{\partial W_{c}}{\partial x} + y \frac{\partial W_{c}}{\partial y} \right) \times \frac{1}{S^{\pm}\pm(z\pm R_{T})}, \\ S^{\pm} &= [(z\pm R_{T})^{2} + x^{2} + y^{2}]^{\frac{1}{2}}, \\ W_{c}(x,y) &= 0.5 \left[1 - \frac{x - x_{0c}}{[(x-x_{0c})^{2} + L_{xc}^{2}]^{\frac{1}{2}}} \right] \times (1 + y^{2}/D_{yc}^{2})^{-1}; \end{split}$$

$$\begin{split} \mathbf{B}_{\mathrm{X}}^{(M)} &= \mathrm{e}^{x/\Delta x} [C_{6}z \cos \psi + (C_{7} + C_{8}y^{2} + C_{9}z^{2}) \sin \psi], \\ \mathbf{B}_{\mathrm{Y}}^{(M)} &= \mathrm{e}^{x/\Delta x} [C_{10}yz \cos \psi + (C_{11}y + C_{12}y^{3} + C_{13}yz^{2}) \sin \psi], \text{ and} \\ \mathbf{B}_{\mathrm{Z}}^{(M)} &= \mathrm{e}^{x/\Delta x} [(C_{14} + C_{15}y^{2} + C_{16}z^{2}) \cos \psi + (C_{17}z + C_{18}zy^{2} + C_{19}z^{3}) \sin \psi], \end{split}$$

where C₆ through C₁₉ are given by:

C _n Kp	= 0,0+	= 1-,1,1+	= 2-,2,2+	= 3 ⁻ ,3,3 ⁺	= 4-,4,4+	≥ 5
C ₆	1.813	2.316	2.641	3.181	3.607	4.090
C ₇	31.10	35.64	42.46	47.50	51.10	49.09
C ₈	-0.07464	-0.0741	-0.07611	-0.1327	-0.1006	-0.0231
C ₉	07764	-0.1081	-0.1579	-0.1864	-0.1927	-0.1359

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C ₁₀	0.003303	0.003924	0.004078	0.01382	0.03353	0.01989
C ₁₁	-1.129	-1.451	-1.391	-1.488	-1.392	-2.298
C ₁₂	0.001663	0.00202	0.00153	0.002962	0.001594	0.004911
C ₁₃	0.000988	0.00111	0.000727	0.000897	0.002439	0.003421
C ₁₄	18.21	21.37	21.86	22.74	22.41	21.79
C ₁₅	-0.03018	-0.04567	-0.04199	-0.04095	-0.04925	-0.05447
C ₁₆	-0.03829	-0.05382	-0.06523	-0.09223	-0.1153	-0.1149
C ₁₇	-0.1283	-0.1457	-0.6412	-1.059	-1.399	-0.2214
C ₁₈	-0.001973	-0.002742	-0.000948	-0.001766	0.000716	-0.01355
C ₁₉	0.000717	0.001244	0.0002276	0.003034	0.002696	0.001185

and where L_y =10.0, D_x =13.0, L_{RC} =5.0, L_T =6.30, γ_T =4.0, δ =0.010, γ_1 =1.0, R_T =30.0, x_{0c} =4.0, L_{xc} =50.0, and D_{yc} =20.0.

Additional parameters relevant to the present invention given in the Tsyganenko reference include (from page 12 of Tsyganenko, *supra*):

Kp parameter	= 0,0+	= 1-,1,1+	= 2-,2,2+	= 3-,3,3+	= 4 ⁻ ,4,4 ⁺	≥ 5-
N	3975	9977	9848	7309	3723	1850
<bc></bc>	15.49	19.06	21.71	25.48	28.58	32.88
C ₁	6.51	8.52	9.75	11.35	12.41	15.12
C ₂	-98.72	-35.64	-77.45	-70.12	-162.50	-128.40
Ċ ₃	-10014	-12800	-14588	-16125	-15806	-16184
C ₄	15.03	14.37	64.85	90.71	160.60	149.10
C ₅	76.62	124.50	123.90	38.08	5.888	215.50

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Δχ	24.74	22.33	20.90	18.64	18.31	19.48
a _{RC}	8.161	8.119	6.283	6.266	6.196	5.831
D_0	2.08	1.664	1.541	0.9351	0.7677	0.3325
γ_{RC}	-0.8799	0.9324	4.183	5.389	5.072	6.472
R _c	9.084	9.238	9.609	8.573	10.06	10.47
G	3.838	2.426	6.591	5.935	6.668	9.081
a _T	13.55	13.81	15.08	15.63	16.11	15.85
D_y	26.94	28.83	30.57	31.47	30.04	25.27
X ₀	5.745	6.052	7.435	8.103	8.260	7.976

It should be noted that the model for B_{TSYG} differs from the model given in Tsyganenko, supra. In the first place, the author has identified several errata to this model. In the second place, the present inventors have modified the model, making C₅ a linear function of Dst for all geomagnetic activity levels. See Boberg et al.

Referring to FIG. 3, the Lorentz equation using a corrected model for the geomagnetic B field is used to calculate the geomagnetic transmission function, using the method depicted in this flowchart. Given the large computational requirements for this calculation, it is preferred to make this calculation in advance for as many orbits as desired, and storing the results as a series of geomagnetic transmission function tables, with the geomagnetic transmission probability given for each particle rigidity (rigidity = momentum/charge), where each table corresponds to the geomagnetic transmission function for a particular orbit. The user may then select from several predefined

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orbits having predefined geomagnetic transmission functions. Alternatively, especially as computer processing power increases, in another embodiment of the invention, this calculation of the geomagnetic transmission function may be performed after a user has specified an orbit.

Referring to 300, the calculation of the geomagnetic transmission function will typically begin by specifying a particle rigidity in GV, a range of particle arrival directions by specifying a range of azimuth angles between θ_{max} and θ_{min} and a range of zenith angles between φ_{max} and φ_{min} , specifying an earth orbit, and specifying a geomagnetic activity level by specifying values for Kp and Dst, where Kp = [IATME or IAGA] planetary Kp index and Dst = the hourly equatorial Dst index. A standard reference for the definition of Kp is Bartels, J., "The standard index, Ks, and the planetary index, Kp", IATME Bulletin 12b, 97 IUGG Pub. Office, Paris, 1949. IATME is the International Association of Terrestrial Magnetism and Electricity. IAGA is the International Association of Geomagnetism and Aeronomy. IUGG is the International Union of Geodesy and Geophysics. A standard reference for the definition of Dst is Sugiura, M., "Hourly values of equatorial Dst for the IGY", Ann. Int. Geophys. Year, 35, 49, 1964. The Dst index was adopted by IAGA at the 1969 meeting in Madrid. IAGA Bulletin 27, 1969, 123, resolution 2. All of the references cited in this paragraph are incorporated herein by reference, in their entireties, for all purposes.

For each value of particle rigidity, the transmission probability will be calculated, and each of these results will be compiled into a table giving the geomagnetic transmission function for a given orbit. A separate geomagnetic transmission function table will be prepared for each specified

15 geomagnetic activity level. The user will also specify an arbitrary number of arrival directions per orbital step M_{adpos}.

Referring to 305, the user will specify an arbitrary number of orbital steps N_{spo} , where N_{spo} is the number of steps in one orbital revolution of the satellite around the Earth's spin axis. The user will also specify a required accuracy for the numerical integration. This embodiment of the invention relies on tracing the particle trajectories backwards from a discrete number of evenly spaced satellite orbital positions. Persons of ordinary skill in the art will recognize, however, that alternative embodiments might be employed, such as embodiments using a Monte Carlo simulation, to trace particle trajectories backwards from randomly selected satellite orbital positions.

Referring to 310, the user will specify the satellite orbit.

Referring to 315, the software will calculate the orbital period from the orbit parameters, using standard orbit generation routines. See Adams et al., NRL Memorandum Report 5099, supra. The software will also calculate a maximum number of steps, and a time per step for the satellite to travel through a step.

Referring to 320, the software will calculate the position of the satellite after each step. See Adams et al., NRL Memorandum Report 5099, supra. The software will also begin tracing the position of a particle incident on the satellite by initializing the particle trajectory time.

Referring to 325, for each of the specified number of arrival directions per orbital step, the software will select a random particle arrival direction. Persons of ordinary skill in the art will

recognize, however, that alternative embodiments might be employed, such as embodiments where a number of regularly spaced arrival directions are used.

Referring to 330, the software will calculate the particle's final position and velocity vector \mathbf{v} from the satellite's latitude, longitude, altitude, and the particle's rigidity, final θ , and final ϕ .

Referring to 330, 340, 345, the software will calculate a new particle position by performing a numerical integration of the Lorentz equation. Although this preferred embodiment uses the Bulirsch-Stoer numerical integration technique, persons of ordinary skill in the art will recognize that other numerical integration techniques are available. This numerical integration is performed on a modified Lorentz equation, where Q is replaced by -Q and t' is replaced by -t'. As persons of ordinary skill in the art will note, the Lorentz equation is invariant if one makes this substitution. This significantly simplifies the numerical integration, because the solution in this case will represent tracing a particle backwards in time from the satellite to the boundary of the magnetosphere (for allowed trajectories) or earth's surface or atmosphere (for forbidden trajectories). This will typically entail far fewer calculations than if particles incident on the magnetosphere were traced forward in time to see if they intersected the satellite. However, persons of ordinary skill in the art will recognize that this tracing forward in time is an equally valid method.

Referring to 350, 355, 360, the software will determine whether, after this iteration of the numerical integration, the particle's new position has placed it within the atmosphere or the solid earth (representing a forbidden trajectory), beyond the magnetosphere as given by the model (representing an allowed trajectory), or still within the magnetosphere (representing an undetermined

trajectory). If the particle is still within the magnetosphere, another iteration of the numerical integration is performed, and a new particle position determined. If the particle is not still in the magnetosphere, i.e., particle has reached either earth's surface or atmosphere (forbidden) or the outside of the magnetosphere (allowed), the software will store this result. This preferred embodiment of the software has an error trapping technique, where if the number of particle steps

Referring to 365, if the software has reached a solution for a given particle arrival direction at a given orbital step, the software will move on to trace the particle arriving at the next arrival direction for that orbital step, if any.

becomes very large, the particle is forced to an allowed or forbidden transition.

Referring to 370, if the software has reached solutions for all the arrival directions for the satellite at a given orbital step, the software moves the satellite along to the next orbital step, if any.

Referring to 375, after all particle tracings for all orbital steps have been performed, the results are compiled as a geomagnetic transmission function of the form $GT(R) = N_{tr}/N_{max}$, where GT is the probability of a particle being transmitted through the magnetosphere, R is particle rigidity, and N_{tr} and N_{max} are a number of particles transmitted and a total number of particles, respectively. Typically, this function will be embodied in a lookup table.

Calculating a geomagnetic transmission function for a given orbit according to the above method is an intensive process. Computing geomagnetic transmission functions for two particular orbits (the proposed space station orbit and a typical space shuttle orbit) required several days of

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processor time on a RISC workstation. Shorter computation cycles could be made on a super-computer, but supercomputer time is in chronic short supply. Accordingly, it is preferred to calculate geomagnetic transmission functions for preselected orbits in advance, and to store these geomagnetic transmission functions for later use by users. This is the approach taken by the inventors in their CREME96 software. In cases where a user specifies an orbit that is not among the precalculated orbits, the CREME96 software will use an earlier model that uses geomagnetic cutoff tables (*see* NRL Memorandum Report 5099, *supra*) rather that geomagnetic transmission functions, and warns the user of this substitution.

Alternatively, a user would specify an orbit, and if the software did not have a precalculated geomagnetic transmission function for this orbit, the software would calculate this geomagnetic transmission function. This embodiment is anticipated to become increasingly desirable as processing capacity increases.

Referring to FIGS 4, 5, 6, and 7, these plots show the geomagnetic transmission function for a common space shuttle orbit (circular orbit, 450 km altitude, 28.5° inclination) under "quiet" (FIG. 4) and "stormy" (FIG. 5) conditions, and the geomagnetic transmission function for the proposed space station orbit (circular orbit, 450 km altitude, 51.6° inclination) under "quiet" (FIG. 6) and "stormy" (FIG. 7) conditions. "Quiet" magnetospheric conditions were taken herein to be Kp = 2 and Dst = -15 nT, while "stormy" magnetospheric conditions were taken herein to be $Kp \ge 5$ and Dst = -300 nT. One sees that (1) transmission drops off sharply with decreasing rigidity, and (2) the differences between stormy and quiet conditions are seen most at lower rigidities.

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The embodiment of the invention implemented in the CREME 96 software operates by prompting a user to specify an orbit and magnetospheric conditions, and then looking up the appropriate geomagnetic transmission function table for the orbit and magnetospheric conditions. The CREME96 software includes lookup tables corresponding to each of FIGS. 4 through 7. This table is then applied to a flux of SEU-inducing particles in the non-trapped energetic particle environment (calculated by another part of the CREME 96 software), to determine the flux of such SEU-inducing particles at the outer surface of the satellite. In other words, the flux from the non-trapped energetic particle environment will be multiplied by the probability (as embodied in the geomagnetic transmission function) that the particles will be transmitted through the magnetosphere, to get the flux at the outer surface of the satellite.

Particle Flux Modeling

Four principal classes of particles are considered to be responsible for most SEUs: galactic cosmic rays (GCRs), anomalous cosmic rays (ACRs), solar protons, and solar heavy ions. See NRL Memorandum Report 4506, supra.

To estimate the effect of these classes of particles on orbiting microelectronics, it is first necessary to estimate the exposure of the microelectronics to these particles. Thus, a feature of any complete model for the effect of cosmic rays on microelectronics will be the modeling of the flux of each of these types of particles.

In the present invention, particle fluxes are determined at the near earth environment. As used herein, the near earth environment refers to the environment found at locations approximately

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1 AU from the sun, but outside of earth's magnetosphere or other localized phenomena that would perturb particle fluxes. After the particle fluxes for each type of particle is determined by the FLUX module of a preferred embodiment of the present invention, other modules may be used to model how the charged particles will induce single event upsets, including what shielding effects (including shielding by the magnetosphere and shielding by the hull of the spacecraft) will reduce this flux at the component level.

In a preferred embodiment of the invention, the flux of GCRs is modeled by the galactic cosmic ray model described by Nymmik et al. See "A Model of Galactic Cosmic Ray Fluxes", Nucl. Tracks Radiat. Meas. 20 427, Nymmik et al. (1992), incorporated herein by reference, in its entirety, for all purposes. Although not essential to the present invention, the authors' updated version of this model is described at http://www.npi.msu.su/gcrf/standart/ISO WD 15390.html, incorporated herein by reference, in its entirety, for all purposes.

In a preferred embodiment of the invention, the flux of ACRs and solar protons are modeled by the model described in Tylka et al, supra.

In a preferred embodiment of the invention, the flux of solar heavy ions is modeled as depicted in FIGS. 8 and 9. This routine depicts how a solar heavy ion flux for a given atomic species at a given kinetic energy, under conditions that are selected from one of three available models: worst day, worst week, and peak flux. As used herein, the worst week solar heavy ion model refers to the model based on particle fluences averaged over a 180 hour span of the solar event beginning at 1300 UT on October 19, 1989. As used herein the worst day solar heavy ion model refers to the model based on particle fluences averaged over an 18 hour span (beginning at 1300 UT on October 24

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20, 1989). As used herein, the peak flux solar heavy ion model refers to the model based on the highest fluxes averaged over 5 minute intervals reported by the Geostationary-orbiting Operational Environmental Satellite (GOES) satellite.

Referring to 800, this routine takes as inputs the atomic number of the selected species, and the kinetic energy for that species whose flux is to be determined. The routine also takes as inputs the user selected baseline model (worst week, worst day, or peak flux).

Referring to 805, if the selected species has atomic number greater than 20, iron is selected as the model spectrum for the flux of this species 810. Otherwise, 815, oxygen is selected as the model spectrum for the flux of this species.

Referring to 820, the next step is to look up an elemental breakpoint EB2. This breakpoint represents a value for the kinetic energy where the relationship between the flux and the kinetic energy changes to a degree where a different mathematical expression for this relationship is warranted. Values for EB2 are given in Table S1:

Table S1: Values for Elemental Breakpoint EB2, in MeV/nuc

Baseline model Elemental spectrum model	worst day or peak flux	worst week
iron	24.23	19.90
oxygen	15.94	12.89

Optionally, this lookup may be performed later in the routine, after the IF step at 825.

Referring to 825, a special case exists for when the baseline model is the worst week model, the model element is Fe, and the kinetic energy being evaluated is above a certain cutoff point

worst week

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 E_{sp} =127.93 MeV/nuc. In this case, referring now to FIG. 9, values for A_{sp} and G_{sp} are looked up from Table S2 and Table S3, respectively, and the unscaled flux is calculated to be $A_{sp} \times En^{-Gsp}$, 900,910.

Table S2: Values for A...:

		Tuble 02. Values for 1	SD*
Flamontol	Baseline model	worst day or peak flux	worst week
Elemental spectrum model			
iron		not applicable	3.16814×10 ⁶
oxygen		not applicable	not applicable

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Table S3: Values for G_{sp} : worst day or peak flux Baseline model Elemental

spectrum model not applicable 2.861 iron

not applicable not applicable oxygen

Referring back to FIG. 8, if En > EB2 (835), values for A_3 and γ_{si} are retrieved from tables S4 and S5, respectively (840), and the unscaled flux is calculated to be $A_3 \times En^{-\gamma si}$ (845).

Table S4: Values for A₃

Elemental spectrum model	Baseline model	worst day or peak flux	worst week	
iron		0.252948×10 ¹⁰	0.249719×10 ⁹	
oxygen		0.106702×10 ¹⁰	0.667628×10°	

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Table S5: Values for γ_{s_1}

Elemental spectrum model	Baseline model	worst day or peak flux	worst week
iron		4.52970	3.7610
oxygen		4.14060	3.76850

Otherwise, for En \leq EB2, values for A₂ and G_{s1} are retrieved from tables S6 and S7, respectively (855), and the unscaled flux is calculated to be A₂exp(-G×En^{1/4})×En^{1/4} (860).

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Table S6: Values for A₂

Elemental spectrum model	Baseline model	worst day or peak flux	worst week
iron		1.8991×10 ⁸	3.0372×10 ⁸
oxygen		4.9518w×10 ⁸	1.1307×10°

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Table S7: Values for G_{s1}

Elemental spectrum model	Baseline model	worst day or peak flux	worst week
iron		5.70	5.70
oxygen		5.70	5.70

For heavier species ($IZ \ge 3$), the flux must be scaled by looking up scale factors for the species being analyzed and the model element (if different from the species being analyzed). These scale factors are taken from Table S8:

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Table S8: Scale factors SF for atomic species with IZ≥3

Atomic Number	Scale Factor	Atomic Number	Scale Fac- tor								
5	0	22	4.377 ×10 ⁻³	39	4.878 ×10 ⁻⁶	56	4.878 ×10 ⁻⁶	73	2.195 ×10 ⁻⁸	90	4.878 ×10 ⁻³
6	4.704 ×10 ⁻¹	23	4.088	40	1 22 ×10 ⁻⁵	57	4.878 ×10 ⁻⁷	74	2.439 ×10 ⁻⁷	91	0
7	1.2059 ×10 ⁻¹	24	1.65 ×10 ⁻²	41	9 756 ×10 ⁻⁷	58	1.22 ×10 ⁻⁶	75	4.878 ×10 ⁻⁸	92	2.927 ×10 ⁻³
8	1	25	5 625 ×10 ⁻³	42	4 878 ×10 ⁻⁶	59	1.951 ×10 ⁻⁷	76	7.317 ×10 ⁻⁷	3	0
9	4 560976 ×10 ⁻⁵	26	1	43	0	60	9 756 ×10 ⁻⁷	77	7.317 ×10 ⁻⁷	4	0
10	2.1312 ×10 ⁻¹	27	1.303 ×10 ⁻²	44	2.195 ×10 ⁻⁶	61	0	78	1.463 ×10 ⁻⁶		
11	1.744715 ×10 ⁻²	28	3.172 ×10 ⁻²	45	4.878 ×10 ⁻⁷	62	2 439 ×10 ⁻⁷	79	2.439 ×10 ⁻⁷		
12	2.0624 ×10 ⁻¹	29	3.048 ×10 ⁻⁴	46	1.463 ×10 ⁻⁶	63	9 756 ×10 ⁻⁸	80	2.439 ×10 ⁻⁷		
13	1.826829 ×10 ⁻²	30	7.457 ×10 ⁻⁴	47	4.878 ×10 ⁻⁷	64	4 878 ×10 ⁻⁷	81	2.195 ×10 ⁻⁷		
14	3 5935 ×10 ⁻¹	31	4.878 ×10 ⁻⁵	48	1.707 ×10 ⁻⁶	65	7.317 ×10 ⁻⁸	82	2.439 ×10 ⁻⁶		
15	2.279675 ×10 ⁻⁴	32	1.22 ×10 ⁻⁴	49	2.195 ×10 ⁻⁷	66	4.878 ×10 ⁻⁷	83	1.463 ×10 ⁻⁷		
16	9.758 ×10 ⁻²	33	7 317 ×10 ⁻⁶	50	4.878 ×10 ⁻⁶	67	9.756 ×10 ⁻⁸	84	0		
17	1.680488 ×10 ⁻⁴	34	7.317 ×10 ⁻⁵	51	3.415 ×10 ⁻⁷	68	2.439 ×10 ⁻⁷	85	0		
18	1.771545 ×10 ⁻³	35	9.756 ×10 ⁻⁶	52	7.317 ×10 ⁻⁶	69	4.878 ×10*	86	0		
19	3.644715 ×10 ⁻⁴	36	4.878 ×10 ⁻⁵	53	1.463 ×10-6	70	1.951 ×10 ⁻⁷	87	0		
20	4.826 ×10 ⁻²	37	7.317 ×10 ⁻⁶	54	6.585 ×10 ⁻⁶	71	4.878 ×10 ⁻⁸	88	0		
21	2.929 ×10 ⁻⁴	38	2.439 ×10 ⁻⁵	55	4.878 ×10 ⁻⁷	72	1.951 ×10 ⁻⁷	89	0		

As embodied in preferred CREME96 software according to the invention, this routine is run for each atomic species in the solar heavy ion flux, and for each value of kinetic energy in the solar

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heavy ion flux. To determine the total solar heavy ion flux, the fluxes for each kinetic energy of each species are added together.

Internet Implementation

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The preferred CREME96 software embodiment of the present invention is large and complex. Moreover, it is anticipated that as models used in the software are improved and/or updated, it will be necessary to perform maintenance on this software. Accordingly, as noted supra, a preferred embodiment of the invention includes a network interface for allowing remote users to access and run the software. Since the internet is a stateless system, a user does not maintain a connection to the server as users in a client-server environment would. One of the goals of the internet implementation of the present invention was to provide users with as much of the functionality of a client-server based system as possible in the stateless system of the internet.

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The preferred system for allowing a large number of users to remotely access and run

software such as CREME96 is a system connected to the internet, or equivalently a system connected

to an intranet. Accordingly, the preferred embodiment of the invention is a server computer (or

equivalently, a plurality of server computers connected in parallel) with the CREME96 software,

with PERL script software (or equivalently Java scripts, or scripts written in some other script

language) for controlling input to and output from the CREME96 software, as well as other

functions, and with listener software and other hardware and software necessary to permit remote

users with internet access to access the CREME96 software through their respective web browsers.

Listings for the preferred software according to the invention are found infra.

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December 31, 1997 (1:24PM)

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Referring to FIG. 10, this figure shows a main menu for the preferred CREME96 software according to the invention. This menu may be accessed through the CREME96 web site maintained at the Naval Research Laboratory having a URL http://crsp3.nrl.navy.mil/creme96/. This page is generated by a PERL script in response to a registered user transmitting to the server a valid username and password. The PERL script will compare the transmitted username and password to a list stored on the server, and if a matching entry is found, the script will generate a CREME96 main menu page customized for that user.

One of the advantages of CREME96 over other web-based applications is that it stores information for each user on the server. This obviates the need for cookies or other downloads to the user's computer. Many users find the downloading of cookies to be objectionable. Moreover, keeping this information on the server may speed up overall operation, because there is no need for the server to download this information to the user, and the user to subsequently upload it to the server. Thus, when a user enters a valid username and password, the script will search a section of the hard drive set aside for that user for any User Request Files (URFs). The user will then be able to pick any of the user's previously stored URFs to use as a data file for a selected routine.

The user has the option, from this main menu, of either running one or more modeling routines, using a selected URF, editing or creating an URF, or running utilities.

The available routines include the GTRN routine for calculating a geomagnetic transmission function as described supra, the FLUX routine for calculating a flux of SEU-inducing particles, as described supra, the TRANS routine for calculating the transmission of SEU-inducing particles through solid shielding, as described supra, the LETSPEC routine for calculating the linear energy 30

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transfer of charged particles to electronic devices, as described supra, and the PUP and HUP routines for calculating the SEU rates for electronic devices exposed to SEU-inducing particles.

The user will select routines to run by clicking a check box next to each of the routines to be run, and picking an URF for each routine to be run from the list of URFs that user has previously stored for that routine. When the user sends this message to the server, by pressing the GO button, a script will generate a system command to run each of these selected routines, using the selected URF as an input file for that routine. As persons of ordinary skill in the art will recognize, a system command is a computer command at the system level. In this case, the system command is to run a program with specified inputs.

Alternatively, if the user selects a radio button next to one of these routines and clicks the GO button, this message from the user will trigger a script to generate and transmit to the user an HTML page with inputs for the data fields in the corresponding URF. If the user has first specified an existing (i.e., stored on the server) URF, the values for the fields in this URF will be included in the generated HTML page, allowing the user to edit the values for the fields in the URF. The URF file name is included in the URF fields, thereby providing the convenience of creating a new URF derived from editing a previous URF, and allowing file storage of both the original and modified URF.

One of the features of the preferred embodiment of the invention is the connectivity between the routines. Referring back to FIG. 1, one sees that the modules of the software are typically executed in the indicated order. In the preferred embodiment, the modules receive input in the form of a URF. The modules also receive input from a file generated by the preceding module (this file 31

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specified by the URF), and/or send output to a file for use by the subsequent module (this file likewise specified by the URF). For instance, the GTRN module will generate a .GT* output file. The FLUX module may use this .GT* file as an input, and will output a .FLX output file. The TRANS module will typically use a .FLX file as an input, and output a .TFX file. The LETSPEC module will typically use a .TFX file (or optionally a .FLX file) as an input, and will output a .LET file. The PUP module will typically use a .TFX file (or optionally a .FLX file) as an input, and will output a .PUP proton-induced upset report file. The HUP module will use a .LET file as an input, and will output a .HUP direct ionization induced upset report file.

Accordingly, by specifying a URF for each of these routines, selecting each of these routines to run, and clicking the GO button, a user will send a message to the server that will cause the scripts to run each of these routines in order, using the outputs from the preceding routines and generating inputs for the subsequent routines. The result of these calculations will be estimates of the single event upset rates for the specified electronics in the specified orbit under the specified conditions. The scripts will generate and transmit an HTML page with these results to the user, who may then use the utilities to download the results, create plots, or perform other functions.

Having described the invention, the following examples are given to illustrate specific applications of the invention, including the best mode now known to perform the invention. These specific examples are not intended to limit the scope of the invention described in this application.

Example 1: CREME96

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Code for CREME96, a preferred embodiment of the present invention, follows. The software may be accessed at its internet site with URL http://crsp3.nrl.navy.mil/creme96/.

Obviously, many modifications and variations of the present invention are possible in light of the above teachings. It is therefore to be understood that, within the scope of the appended claims, the invention may be practiced otherwise than as specifically described.

CLAIMS

What is claimed is:

- 1. A method for determining a geomagnetic transmission function for the transmission of a population of particles, said particles having one or more rigidities, to a satellite in a known earth orbit, under known geomagnetic conditions, comprising the steps:
- (A) dividing said orbit into a plurality of sequential steps comprising at least a first step and a last step, each sequential pair of said steps being connected at a point, each of said points corresponding to a position of said satellite along said orbit, and specifying a value for Kp and a value for Dst to specify said geomagnetic conditions;
- (B) for a plurality of said points, selecting a plurality of arrival directions for the arrival of said particles at said satellites, wherein each of said arrival directions represents either an allowed or a forbidden trajectory of one of said particles; and
- (C) for each of said arrival directions at each of said plurality of said points, determining whether said arrival direction represents an allowed or a forbidden trajectory by tracing a path of said particle to arrive at said point with said arrival direction, until said particle path intersects a boundary of earth's magnetosphere, thereby indicating an allowed trajectory, or until said particle path intersects either earth's atmosphere or earth's surface, thereby indicating a forbidden trajectory, wherein said tracing of said path is performed by integrating in the time domain the Lorentz equation

$$\mathbf{F} = m \gamma \frac{d \mathbf{v}}{d(-t')} = -Q \mathbf{v} \times \mathbf{B}$$

wherein **F** is the force vector acting on said particle, m is mass, γ is $1/(1-\mathbf{v}^2/c^2)^{-1/2}$, **v** is said particle's velocity vector, t' is the travel time of said particle, Q is the charge of said particle, and **B** is the magnetic field vector acting on said particle, wherein **B** is given by

$$\mathbf{B} = \mathbf{B}_{\text{IGRF}}(\mathbf{r}, t') + \mathbf{B}_{\text{TSYG}}(\mathbf{K}\mathbf{p}, \mathbf{r}, t')$$

wherein \mathbf{B}_{IGRF} is the International Geomagnetic Reference Field promulgated by the International Association of Geomagnetism and Aeronomy, and \mathbf{B}_{TSYG} is the modified Tsyganenko field given by the sum of $\mathbf{B}_{Xmg}^{(T)} + \mathbf{B}_{Ymg}^{(T)} + \mathbf{B}_{Zmg}^{(T)} + \mathbf{B}_{Zmg}^{(RC)} + \mathbf{B}_{Ymg}^{(RC)} + \mathbf{B}_{Zmg}^{(RC)} + \mathbf{B}_{Zmg}^{(RC)} + \mathbf{B}_{Ysm}^{(C)} + \mathbf{B}_{Zsm}^{(C)} +$

$$\mathbf{B}_{\mathbf{X}}^{(T)} = \mathbf{Q}_{\mathbf{t}} \times \mathbf{z}_{\mathbf{r}};$$

$$\mathbf{B}_{\mathbf{Y}}^{(T)} = \mathbf{Q}_{t} \mathbf{y} \mathbf{z}_{r};$$

$$\mathbf{B}_{z}(T) = \frac{\mathbf{W}(\mathbf{x}, \mathbf{y})}{\mathbf{S}_{T}} \left(\mathbf{C}_{1} + \mathbf{C}_{2} \frac{\mathbf{a}_{T} + \boldsymbol{\xi}_{T}}{\mathbf{S}_{T}^{2}} \right) + \frac{\mathbf{x} \frac{\partial \mathbf{W}}{\partial \mathbf{x}} + \mathbf{y} \frac{\partial \mathbf{W}}{\partial \mathbf{y}}}{\mathbf{S}_{T} + \mathbf{a}_{T} + \boldsymbol{\xi}_{T}} \times \left(\mathbf{C}_{1} + \frac{\mathbf{C}_{2}}{\mathbf{S}_{T}} \right) + \mathbf{B}_{x}(T) \frac{\partial \mathbf{z}_{s}}{\partial \mathbf{x}} + \mathbf{B}_{y}(T) \frac{\partial \mathbf{z}_{s}}{\partial \mathbf{y}} - \mathbf{Q}_{T} \mathbf{D}_{T} \left(\mathbf{x} \frac{\partial \mathbf{D}_{T}}{\partial \mathbf{x}} + \mathbf{y} \frac{\partial \mathbf{D}_{T}}{\partial \mathbf{y}} \right)$$

wherein W(x,y) = 0.5
$$\left(1 - \frac{x - x_0}{\left[\left(x - x_0\right)^2 + D_x^2\right]^{1/2}}\right) \times \left(1 + \frac{y^2}{D_y^2}\right)^{-1}$$

wherein
$$Q_T = \frac{W(x,y)}{\xi_T S_T} \left[\frac{C_1}{S_T + a_T + x_T} + \frac{C_2}{S_T^2} \right];$$

wherein $z_r = z - z_s(x, y, \psi)$,

$$\begin{split} z_{\mathrm{s}}(\mathbf{x},\mathbf{y},\psi) &= 0.5\tan\psi \Big(\mathbf{x} + \mathbf{R}_{\mathrm{c}} - \sqrt{\left(\mathbf{x} + \mathbf{R}_{\mathrm{c}}\right)^2 + 16}\Big) \\ &- \mathrm{Gsin}\,\psi \cdot \mathbf{y}^4 \Big(\mathbf{y}^4 + \mathbf{L}_{\mathrm{y}}^4\Big)^{-1}, \\ \mathrm{wherein} \qquad \mathbf{S}_{\mathrm{T,RC}} &= \sqrt{\rho^2 + \left(\mathbf{a}_{\mathrm{T,RC}} + \xi_{\mathrm{T,RC}}\right)^2}, \\ \xi_{\mathrm{T,RC}} &= \sqrt{\mathbf{z}_{\mathrm{r}}^2 + \mathbf{D}_{\mathrm{T,RC}}^2}, \\ D_T &= D_0 + \delta \mathbf{y}^2 + \gamma_T h_T(\mathbf{x}) + \gamma_1 h_1(\mathbf{x}), \end{split}$$

wherein
$$C_1 = -98.72$$
 when $Kp = 0$, 0^+ , $C_1 = -35.64$ when $Kp = 1^-$, 1 , 1^+ , $C_1 = -77.45$ when $Kp = 2^-$, 2 , 2^+ , 1 when 1

 $C_1 = -128.4$ when $Kp \ge 5^-$,

wherein $C_2 = -10014$ when Kp = 0, 0^+ , $C_2 = -12800$ when $Kp = 1^-, 1, 1^+$, $C_2 = -14588$ when $Kp = 2^-, 2, 2^+$, $C_2 = -16125$ when $Kp = 3^-, 3, 3^+$, $C_2 = -15806$ when $Kp = 4^-, 4, 4^+$, $C_2 = -16184$ when $Kp \ge 5^-$,

wherein $a_T = 13.55$ when Kp = 0, 0^+ , $a_T = 13.81$ when $Kp = 1^-$, 1, 1^+ , $a_T = 15.08$ when $Kp = 2^-$, 2, 2^+ , $a_T = 15.63$ when $Kp = 3^-$, 3, 3^+ , $a_T = 16.11$ when $Kp = 4^-$, 4, 4^+ , $a_T = 15.85$ when $Kp \ge 5^-$,

wherein $D_0 = 2.08$ when Kp = 0, 0^+ , $D_0 = 1.664$ when $Kp = 1^-$, 1, 1^+ , $D_0 = 1.541$ when $Kp = 2^-$, 2, 2^+ , $D_0 = 0.9351$ when $Kp = 3^-$, 3, 3^+ , $D_0 = 0.7677$ when $Kp = 4^-$, 4, 4^+ ,

wherein $R_c = 9.084$ when Kp = 0, 0^+ , $R_c = 9.238$ when $Kp = 1^-,1,1^+$, $R_c = 9.609$, when $Kp = 2^-,2,2^+$, $R_c = 8.573$ when $Kp = 3^-,3,3^+$, $R_c = 10.06$ when $Kp = 4^-,4,4^+$, $R_c = 10.47$ when $Kp \ge 5^-$,

wherein $L_y = 10 R_E$,

$$\mathbf{B}_{\mathbf{X}}^{(\mathrm{RC})} = \mathbf{Q}_{\mathrm{RC}} \mathbf{X} \mathbf{z}_{\mathrm{r}};$$

$$\mathbf{B}_{\mathbf{Y}}^{(\mathbf{RC})} = \mathbf{Q}_{\mathbf{RC}} \mathbf{y} \mathbf{z}_{\mathbf{r}};$$

$$\mathbf{B}_{\mathrm{Z}}^{(\mathrm{RC})} = C_{5} \frac{2(a_{\mathrm{RC}} + \xi_{\mathrm{RC}})^{2} - \rho^{2}}{S_{\mathrm{RC}}^{5}} + B_{\mathrm{X}}^{\mathrm{RC}} \frac{\partial z_{\mathrm{s}}}{\partial x} + B_{\mathrm{Y}}^{\mathrm{RC}} \frac{\partial z_{\mathrm{s}}}{\partial y} - Q_{\mathrm{RC}} D_{\mathrm{RC}} x \frac{\partial D_{\mathrm{RC}}}{\partial x};$$

wherein
$$Q_{RC} = 3C_5 \xi_{RC}^{-1} S_{RC}^{-5} (a_{RC} + \xi_{RC})$$

$$\begin{split} &D_{RC} = D_0 + \gamma_{RC} h_{RC}(x) + \gamma_1 h_1(x) \\ &h_{T,RC} = 0.5 [1 + x(x^2 + L_{T,RC}^2)^{-1/2}], \\ &h_1 = 0.5 \{1 - (x + 16)[(x + 16)^2 + 36]^{-1/2}\}, \end{split}$$

$$\begin{split} C_{5}(Dst) &= -10220 + 408.5 \cdot Dst \\ \mathbf{B}_{XYZ}^{(C)} &= C_{3}(F_{x,y,z}^{+} + F_{x,y,z}^{-}) + C_{4}(F_{x,y,z}^{+} - F_{x,y,z}^{-}), \text{ wherein} \\ &\left\{\frac{F_{x}^{\pm}}{F_{y}^{\pm}}\right\} = \pm \frac{W_{c}(x,y)}{S^{\pm}[S^{\pm}\pm(z\pm R_{T})]} \times \left\{\frac{x}{y}\right\}, \\ F_{z}^{\pm} &= \frac{W_{c}(x,y)}{S^{\pm}} + \left(x\frac{\partial W_{c}}{\partial x} + y\frac{\partial W_{c}}{\partial y}\right) \times \frac{1}{S^{\pm}\pm(z\pm R_{T})}, \\ S^{\pm} &= [(z\pm R_{T})^{2} + x^{2} + y^{2}]^{\frac{1}{2}}, \\ W_{c}(x,y) &= 0.5 \left[1 - \frac{x - x_{0c}}{[(x-x_{0c})^{2} + L_{yc}^{2}]^{\frac{1}{2}}}\right] \times (1 + y^{2}/D_{yc}^{2})^{-1}; \end{split}$$

$$\begin{split} &\mathbf{B}_{\mathrm{X}}^{(\mathrm{M})} = e^{x/\Delta x} [C_{6}z\cos\psi + (C_{7} + C_{8}y^{2} + C_{9}z^{2})\sin\psi], \\ &\mathbf{B}_{\mathrm{Y}}^{(\mathrm{M})} = e^{x/\Delta x} [C_{10}yz\cos\psi + (C_{11}y + C_{12}y^{3} + C_{13}yz^{2})\sin\psi], \text{ and} \\ &\mathbf{B}_{\mathrm{Z}}^{(\mathrm{M})} = e^{x/\Delta x} [(C_{14} + C_{15}y^{2} + C_{16}z^{2})\cos\psi + (C_{17}z + C_{18}zy^{2} + C_{19}z^{3})\sin\psi], \end{split}$$

wherein C_6 through C_{19} are given by:

C_n Kp	$=0,0^{+}$	= 1-,1,1+	= 2~,2,2+	= 3 ⁻ ,3,3 ⁺	= 4 ⁻ ,4,4 ⁺	≥ 5	
C_6	1.813	2.316	2.641	3.181	3.607	4.090	
C ₇	31.10	35.64	42.46	47.50	51.10	49.09	
C ₈	-0.07464	-0.0741	0.0741 -0.07611		-0.1006	-0.0231	
C ₉	07764	-0.1081	-0.1579	-0.1864	-0.1927	-0.1359	
C ₁₀	0.003303	0.003924	0.004078	0.01382	0.03353	0.01989	
C ₁₁	-1.129	-1.451	-1.391	-1.488	-1.392	-2.298	
C ₁₂	0.001663	0.00202	0.00153	0.002962	0.001594	0.004911	
C ₁₃	0.000988	0.00111	0.000727	0.000897	0.002439	0.003421	
C ₁₄	18.21	21.37	21.86	22.74	22.41	21.79	
C ₁₅	-0.03018	-0.04567	-0.04199	-0.04095	-0.04925	-0.05447	
C ₁₆	-0.03829	-0.05382	-0.06523	-0.09223	-0.1153	-0.1149	
C ₁₇	-0.1283	-0.1457	-0.6412	-1.059	-1.399	-0.2214	
C ₁₈	-0.001973	-0.002742	-0.000948	-0.001766	0.000716	-0.01355	
C ₁₉	0.000717	0.001244	0.0002276	0.003034	0.002696	0.001185	

and wherein L_y =10.0, D_x =13.0, L_{RC} =5.0, L_T =6.30, γ_T =4.0, δ =0.010, γ_1 =1.0, R_T =30.0, x_{0c} =4.0, L_{xc} ²=50.0, and D_{yc} =20.0,

and thereby determining whether said particle's trajectory intersects either the boundary of earth's magnetosphere, thereby indicating an allowed trajectory, or intersecting earth's surface or earth's atmosphere, thereby indicating a forbidden trajectory.

2. The method of claim 1, wherein said plurality of arrival directions comprises one or more randomly or pseudorandomly selected arrival directions.

- 3. The method of claim 1, further comprising the step:
- (D) for each particle rigidity, determine what fraction of particles of that rigidity will be transmitted to said satellite.
- 4. The method of claim 1, wherein said plurality of points comprises a complete set of points for said orbit.
- 5. The method of claim 1, wherein said plurality of points comprises a set of points for a portion of said orbit.
- 6. A method for determining, for a given particle environment outside of earth's magnetosphere, what portion of a population of particles having one or more rigidities making up said particle environment will be transmitted to a satellite in a known earth orbit, comprising the steps:
- (A) performing steps (A) through (C) of claim 1, thereby computing a geomagnetic transmission function for said population of particles; and
- (B) applying said geomagnetic transmission function to said population of particles, thereby determining what portion of that population of particles will be transmitted to said satellite.
- 7. A method for determining, for a given particle environment outside of earth's magnetosphere, what portion of a population of particles having one or more rigidities making up said particle environment will be transmitted to a satellite in earth orbit, comprising the steps:
 - (A) prompting a user to specify an earth orbit;

- (B) determining whether said orbit is among a group of preselected orbits, each of said orbits in said preselected group of orbits having an associated predetermined geomagnetic transmission function for a range of particle rigidities, each of said predetermined geomagnetic transmission functions having been prepared in accordance with claim 1; and
- (C) for the case wherein said orbit is among said preselected group of orbits, applying said predetermined geomagnetic transmission function for said orbit to said particle environment outside earth's magnetosphere.
- 8. The method of claim 7, wherein said preselected group of orbits comprises a quiet shuttle orbit having a 450 km altitude and a 28.5° inclination, a disturbed shuttle orbit having a 450 km altitude and a 28.5° inclination, a quiet space station orbit having a 450 km altitude and a 51.6° inclination, and a disturbed space station orbit having a 450 km altitude and a 51.6° inclination, wherein said quiet shuttle orbit has a geomagnetic transmission function given by FIG. 4, wherein said disturbed shuttle orbit has a geomagnetic transmission function given by FIG. 5, wherein said quiet space station orbit has a geomagnetic transmission function given by FIG. 6, and wherein said disturbed space station orbit has a geomagnetic transmission function given by FIG. 7.
- 9. The method of claim 7, further comprising the step:
- (D) for the case wherein said orbit is not among said preselected group of orbits, performing a step selected from the group consisting of (a) returning an error message to a user, and (b) computing a geomagnetic transmission function for said orbit, in accordance with the method of claim 1.

- 10. A method for determining a flux of a species of solar ions having a specified atomic number between 3 and 92, and a specified kinetic energy, for a satellite in a near earth orbit, comprising the steps:
 - (A) specifying an atomic number for solar ions for evaluation;
 - (B) specifying a kinetic energy for said solar ions;
- (C) specifying a baseline model for said flux of solar ions, wherein said baseline model is selected from the group consisting of a worst day model, a worst week model, and a peak flux model;
- (D) in the case wherein said specified atomic number is greater than 20, selecting iron as an elemental spectrum model;
- (E) in the case wherein said specified atomic number is less than or equal to 20, selecting oxygen as an elemental spectrum model;
- (F) looking up a value for an elemental breakpoint, wherein said elemental breakpoint is a function of said elemental spectrum model and said baseline model, and wherein said elemental breakpoint is selected from the table:

Elemental spectrum model	Baseline model	worst day or peak flux	worst week
iron		24.23 MeV/nuc	19.90 MeV/nuc
oxygen		15.94 MeV/nuc	12.89 MeV/nuc

(G) in the case wherein said baseline model is said worst week model, and said elemental spectrum model is iron, and said kinetic energy is greater than 127.93 MeV/nuc, calculating an unscaled flux, wherein said unscaled flux equals $A_{sp} \times \left(En/\frac{\text{MeV}_{nuc}}{\text{nuc}}\right)^{-Gsp}$, wherein $A_{sp} = 3.16814 \times 10^6$

(cm² sr MeV/nuc)⁻¹, $\left(En/\frac{\text{MeV}}{\text{nuc}}\right)$ is said kinetic energy, normalized to be dimensionless, and G_{sp} = 2.861;

(H) in the case wherein the condition recited in step (G) is not satisfied, and wherein said kinetic energy is greater than said elemental breakpoint, calculating an unscaled flux wherein said unscaled flux equals $A_3 \times EN^{\gamma_{SI}}$, wherein A_3 is a function of said elemental spectrum model and said baseline model, and wherein said A_3 is selected from the table:

Baseline r. Elemental spectrum model	model worst day or peak flux in (cm² sr MeV/nuc)-1	worst week in (cm ² sr MeV/nuc) ⁻¹
iron	0.252948×10 ¹⁰	0.249719×10°
oxygen	0.106702×10 ¹⁰	0.667628×10°

and wherein γ_{s_1} is a spectral index and is a function of said elemental spectrum model and said baseline model, and wherein said γ_{s_1} is selected from the table:

Baseline Elemental spectrum model	model worst day or peak flux	worst week	
iron	-4.52970	-3.7610	
oxygen	-4.14060	-3.76850	

(I) in the case wherein the condition recited in step (G) is not satisfied, and wherein said kinetic energy is less than or equal to said elemental breakpoint, calculating an unscaled flux wherein said unscaled flux equals $A_2 \exp(-G \times En^{1/4}) \times En^{1/4}$, wherein A_2 is a function of said elemental spectrum model and said baseline model, and wherein said A_2 is selected from the table:

Baseline model Elemental spectrum model	worst day or peak flux in (cm ² sr MeV/nuc) ⁻¹	worst week in (cm ² sr MeV/nuc) ⁻¹
---	--	---

iron	1.8991×10 ⁸	3.0372×10 ⁸			
oxygen	4.9518×10 ⁸	1.1307×10 ⁹			

and wherein G_{si} is a spectral index and is a function of said elemental spectrum model and said baseline model, and wherein said G_{si} is selected from the table:

Elemental spectrum model	Baseline model	worst day or peak flux	worst week
iron		5.70	5.70
oxygen		5.70	5.70

[;] and

(J) in the case where said atomic number is between 3 and 92, inclusive, calculating a solar ion flux for said specified atomic number and said specified kinetic energy by multiplying said unscaled flux by a scale factor ratio, said scale factor ratio being the ratio of a scale factor for an element having said selected atomic number over a scale factor for said spectrum model element, wherein said scale factors are selected from the table:

Atomic Number	Scale Factor	Atomic Number	Scale Factor	Atomic Number	Scale Factor	Atomic Number	Scale	Atomic	Scale	Atomic	Scale
ranibei		Number	Factor	Number	ractor	Number	Factor	Number	Factor	Number	Factor
5	0	22	4 377 ×10 ⁻³	39	4 878 ×10 ⁻⁶	56	4 878 ×10 ⁻⁶	73	2.195 ×10 ⁻⁸	90	4.878 ×10 ⁻⁸
6	4 704 ×10 ⁻¹	23	4.088	40	1.22 ×10 ⁻⁵	57	4.878 ×10 ⁻⁷	74	2.439 ×10 ⁻⁷	91	0 -
7	1.2059 ×10 ⁻¹	24	1.65 ×10 ⁻²	41	9.756 ×10 ⁻⁷	58	1.22 ×10 ⁻⁶	75	4.878 ×10 ⁻⁸	92	2.927 ×10 ⁻⁸
8	1	25	5 625 ×10 ⁻³	42	4.878 ×10 ⁻⁶	59	1 951 ×10 ⁻⁷	76	7.317 ×10 ⁻⁷	3	0
9	4.560976 ×10 ⁻⁵	26	1	43	0	60	9 756 ×10 ⁻⁷	77	7.317 ×10 ⁻⁷	4	0
10	2.1312 ×10 ⁻¹	27	1.303 ×10 ⁻²	44	2.195 ×10 ⁻⁶	61	0	78	1.463 ×10 ⁻⁶		
11	1.744715 ×10 ⁻²	28	3.172 ×10 ⁻²	45	4 878 ×10 ⁻⁷	62	2.439 ×10 ⁻⁷	79	2.439 ×10 ⁻⁷		
12	2.0624 ×10 ⁻¹	29	3.048 ×10 ⁻⁴	46	1.463 ×10 ⁻⁶	63	9 756 ×10 ⁻⁸	80	2.439 ×10 ⁻⁷		
13	1.826829 ×10 ⁻²	30	7.457 ×10 ⁻⁴	47	4 878 ×10 ⁻⁷	64	4 878 ×10 ⁻⁷	81	2.195 ×10 ⁻⁷		
14	3 5935 ×10 ⁻¹	31	4.878 ×10 ⁻⁵	48	1.707 ×10 ⁻⁶	65	7 317 ×10 ⁻⁸	82	2.439 ×10 ⁻⁶		
15	2.279675 ×10 ⁻⁴	32	1 22 ×10 ⁻⁴	49	2 195 ×10 ⁻⁷	66	4.878 ×10 ⁻⁷	83	1.463 ×10 ⁻⁷		
16	9.758 ×10 ⁻²	33	7.317 ×10 ⁻⁶	50	4.878 ×10 ⁻⁶	67	9.756 ×10 ⁻⁸	84	0		
17	1 680488 ×10 ⁻⁴	34	7 317 ×10 ⁻⁵	51	3.415 ×10 ⁻⁷	68	2 439 ×10 ⁻⁷	85	0		
18	1 771545 ×10 ⁻³	35	9.756 ×10 ⁻⁶	52	7 317 ×10 ⁻⁶	69	4 878 ×10 ⁻⁸	86	0		
19	3.644715 ×10 ⁻⁴	36	4.878 ×10 ⁻⁵	53	1.463 ×10 ⁻⁶	70	1.951 ×10 ⁻⁷	87	0		
20	4.826 ×10 ⁻²	37	7.317 ×10 ⁻⁶	54	6.585 ×10 ⁻⁶	71	4.878 ×10 ⁻⁸	88	0		
21	2.929 ×10 ⁻⁴	38	2.439 ×10 ⁻⁵	55	4.878 ×10 ⁻⁷	72	1.951 ×10 ⁻⁷	89	0		

wherein elements having atomic number of 20 or less are scaled to oxygen, and elements having higher atomic numbers are scaled to iron.

- 11. A method for determining a flux of solar ions of species having a range of atomic numbers between 3 and 92, wherein each of said species includes particles having a range of specified kinetic energies, for a satellite in a near earth orbit, comprising the steps:
- (A) for each kinetic energy of each of said species, performing steps (A) through (K) of claim 10, and storing a flux value for each of said kinetic energies for each of said species; and
 - (B) adding each of said flux values to obtain a total solar ion flux value.
- 12. A method for modeling the effects of cosmic rays on microelectronics on a computer connected to the internet, comprising the steps:
- (A) receiving a login message from a remote user connected to the internet, and generating and transmitting back to said user with a script in response thereto an HTML main menu page, wherein said main menu page comprises prompts for said user to run one or more routines selected from the group consisting of: (1) a routine for calculating a geomagnetic transmission function for SEU-inducing particles, (2) a routine for calculating a flux of SEU-inducing particles in the near earth environment or in the environment shielded by earth's magnetosphere, (3) a routine for calculating a solid shielding transport function for SEU-inducing particles, (4) a routine for calculating a proton induced single event upset rate, (5) a routine for calculating a linear energy transfer rate, and (6) a routine for calculating a heavy ion induced single event upset rate, said main menu page further comprises prompts for said user to select a user request file for each of said routines;
- (B) in response to a message from said user to run at least one of said routines, using a user request file specified by said user, generating and executing with a script a system command for said

computer to run each of said specified routines; and

- (C) in response to the completion of all of said routines specified by said user, generating and transmitting to said user with a script an HTML page notifying said user of said completion.
- 13. The method of claim 12, wherein said main menu page generated and transmitted to said user further comprises prompts for said user to create or edit a user request file for any one of said routines, in lieu of running said one or more of said routines, and further comprising the step:
- (D) in response to a message from said user to create or edit a user request file, generating and transmitting to said user with a script an HTML page with prompts for the entry of fields for said user request file; and
- (E) in response to said user transmitting information for said user request file, storing said information in a user request file with a script.
- 14. The method of claim 12, wherein said main menu page comprises prompts for said user to run a routine for calculating a geomagnetic transmission function for SEU-inducing particles, and wherein said step of generating and executing with a script a system command for said computer to run each of said specified routines comprises at least generating and executing a system command for said computer to run said routine for calculating a geomagnetic transmission function for SEU-inducing particles, in accordance with claim 1
- 15. The method of claim 12, wherein said main menu page comprises prompts for said user to run a routine for calculating a flux of SEU-inducing particles in the near earth environment or in the

environment shielded by earth's magnetosphere, and wherein said step of generating and executing with a script a system command for said computer to run each of said specified routines comprises at least generating and executing a system command for said computer to run said routine for calculating said flux of SEU-inducing particles in the near earth environment or in the environment shielded by earth's magnetosphere, wherein said routine comprises calculating a flux of solar heavy ions in accordance with claim 10.

ABSTRACT OF THE DISCLOSURE

An aspect of the present invention is a method and apparatus for computing a geomagnetic transmission function. This apparatus includes a programmed digital computer running modeling software for modeling the transmission of cosmic ray particles through the magnetosphere. The software includes a model representing a solution to the Lorentz equation in a magnetic field given by $\mathbf{B} = \mathbf{B}_{\text{IGRF}}(\mathbf{r}, t') + \mathbf{B}_{\text{TSYG}}(\mathbf{Kp}, \mathbf{r}, t')$. Another aspect of the present invention is a method and apparatus for computing a flux of particles at the outer surface of a satellite comprising, inter alia, an improved method and apparatus for computing a flux of solar heavy ions. This apparatus includes a programmed digital computer running modeling software for modeling the flux of cosmic ray particles through the outer surface of a satellite. Another aspect of the invention is a method and apparatus comprising a programmed digital computer running modeling software for modeling the effect of cosmic rays on microelectronics, where this software embodies at least one of the two foregoing aspects of the invention. Another aspect of the invention is a preferred embodiment of a method and apparatus comprising a programmed digital computer running modeling software for modeling the effect of cosmic rays on microelectronics, where this software embodies at least one of the two foregoing aspects of the invention, where this preferred embodiment is connected to a network, typically the internet, to permit remote users to use the invention.

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REAL FUNCTION ACR FLUX(IZ, IQ, EN, YEAR)

Returns the interplanetary anomalous cosmic ray (ACR) flux for element IZ in charge state IQ at energy EN (in MeV/nuc) for year YEAR (ie, decimal year 1987.23)

Flux returned in unit of ions/m2-s-sr-MeV/nuc

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This routine based on results from several sources:

(1) ACR Elemental Composition

Cummings & Stone discovered that all ACR spectra could be mapped to a common spectral shape by using flux and energy scaling factors, contained here in the arrays FFAC and EFAC, respectively. The values of these scaling factors are taken from various sources. Unless otherwise noted, they are given in:

Cummings & Stone, Proc. 20th ICRC (Moscow), 3, 413-416 (1987).

FFAC values for N, Ne modified to match the interplanetary abundances reported from SAMPEX at 16-25 MeV/nuc: Selesnick et al., JGR 100, 9503-9518 (1995).

He spectrum taken from:

Cummings, Stone, & Webber, Ap.J. Lett. 287, L199-L103 (1984), which is based on Voyager observations in 9/1977-2/1978, when the spacecraft were still near Earth ($\sim 1.3-2.5$ AU)

Relative abundance and spectrum of ACR H at 1 AU estimated by: Mewaldt, Proc. 24th ICRC (Rome) 4, 808-811 (1995). (The spectrum here roughly matches this.)

(2) Spectral shape and distribution of charge states:

Based on SAMPEX observations and modeling results presented by R.A. Mewaldt et al., Astrophysical Journal Letters 466, L43-L46 (1996).

(3) Temporal variation and solar-cycle modulation:

Based on the timeline of 8-27 MeV/nuc oxygen for 1968-1994 as reported by Mewaldt et al. GRL 20, 2263-2266 (1993) and augmented during the 1985-95 time period by Cosmos measurements reported by: Beaujean et al. Proc. 24th ICRC (Rome) 4, 832-835 (1994).

IMPLICIT NONE

INTEGER*4 IZ, IQ, NELM

PARAMETER (NELM=18)

REAL*4 EN, YEAR, FPEAK, AMASS, FFAC, EFAC, EN1

DIMENSION FPEAK(4), FFAC(NELM), EFAC(NELM), AMASS(NELM)

DATA FPEAK/3.935,0.3808,0.1014,3.01E-2/

DATA FFAC/3.6,4.9,3*0.0,0.0075,0.127,1.0,0.0,0.40,7*0.0,0.019/

```
DATA EFAC/9.0,2.6,3*1.0,1.38,1.14,1.0,1.0,0.64,7*1.0,0.36/
         DATA AMASS/
       & 1.00794,4.002602,6.941,9.012182,10.811,12.011,14.00674,15.9994,
       £ 18.9984032,20.1797,22.989768,24.305,26.981539,28.0855,30.973762,
       & 32.066,35.4527,39.948/
        REAL*4 AO, Q, AN, FP, BETA, ANORM, EPEAK
        DATA A0/1.70/
        REAL*4 SMNORM/0.5534E+4/
        REAL*4 ACRO_TIMELINE
 C
        ACR FLUX=0.0
        IF (IZ.GT.NELM) RETURN
        IF (FFAC(IZ).LE.0) RETURN
        IF (EN.LT.1.0 .or. EN.GT.1000.) RETURN
        IF (IQ.GT.IZ) RETURN
        Q=IQ*1.0
        AN=AMASS(IZ)
        IF (IQ.LE.4) THEN
            FP=FPEAK(IQ)
        ELSEIF (IQ.GT.4) THEN
            FP=3.935/Q**3.52
        ENDIF
        EPEAK=6.73*Q**0.91
        FP=FP*FFAC(IZ)
        BETA=1.0/A0/EPEAK**A0
        ANORM=SMNORM*FP/EPEAK/EXP(-BETA*EPEAK**A0)
        EN1=EN/EFAC(IZ)
        ACR_FLUX=ANORM*EN1*EXP(-BETA*EN1**A0)
        Solar modulation factor:
       ACR_FLUX=ACR FLUX*ACRO TIMELINE (YEAR)
       RETURN
        END
        REAL FUNCTION ACRO TIMELINE (USERX)
        Function to model the solar cycle variation of the anomalous
C
        component at 1 AU. Based on the timeline of 8-27 MeV/nuc oxygen
C
        for 1968-1994 as reported by Mewaldt et al. GRL 20, 2263-2266 (1993)
С
        and augmented during the 1985-95 time period by Cosmos measurements
C
        reported by: Beaujean et al. Proc. 24th ICRC (Rome) 4, 832-835 (1994).
C
C Calculates a y-value, along a line drawn through data points, for a given
C year (USERX) on Fig.3 of Mewaldt et al.
        IMPLICIT NONE
C Argument declarations
        REAL USERX
C Local declarations
        INTEGER KMAX, I
        PARAMETER (KMAX=5)
        REAL XVAL(KMAX+1), YVAL(KMAX+1), SLOPE(KMAX), YEAR
        DATA XVAL /1.9685E+03,1.9715E+03,1.9778E+03,1.9825E+03,
     & 1.9873E+03,1.9903E+03/
        DATA YVAL /5.8651E-09,1.2123E-06,2.0350E-06,6.4463E-09,
     & 1.7630E-06,5.8651E-09/
        DATA SLOPE /1.782245,8.206459E-02,-1.217980,1.177813,-1.896111/
```

```
C
          DATA XVAL /1.9685E+03,1.9717E+03,1.9778E+03,1.9825E+03,
С
       & 1.9873E+03,1.9903E+03/
C
         DATA YVAL /5.8651E-09,2.035E-06,2.0350E-06,6.4463E-09.
С
      & 1.7630E-06,5.8651E-09/
C
         DATA SLOPE /1.782245,0.00,-1.217980,1.177813,-1.896111/
     &
C
      Evaluate which slope to use and calculate y-value
      CALL ACRO YEAR (USERX, XVAL(1), XVAL(KMAX+1), YEAR)
      DO I=1, KMAX
          IF ((YEAR.GE.XVAL(I)).AND.(YEAR.LT.XVAL(I+1)))
             ACRO_TIMELINE=EXP(ALOG(YVAL(I+1))-
     ٠.
     &
                           (SLOPE(I) * (XVAL(I+1) - YEAR)))
      ENDDO
      RETURN
      END
      SUBROUTINE ACRO_YEAR (USERX, LOWERX, UPPERX, YEAR)
C Evaluates a given year to see if it falls within the range of 1967.9-1990.5.
C If it doesn't, USERX is updated by either adding or subtracting a factor
C of 21.8 so that it does fall within the specified range.
C Declarations
      IMPLICIT NONE
      REAL USERX, LOWERX, UPPERX, DIFF, RMDR, YEAR
      INTEGER FACTOR
C Evaluate and modify USERX
      IF (USERX .LT. LOWERX) THEN
         DIFF=LOWERX-USERX
         RMDR=DIFF/21.8
         FACTOR=INT(RMDR)+1
         YEAR=USERX+REAL(FACTOR)*21.8
      ELSEIF (USERX .GT. UPPERX) THEN
         DIFF=USERX-UPPERX
         RMDR=DIFF/21.8
         FACTOR=INT (RMDR) +1
         YEAR=USERX-REAL (FACTOR) *21.8
      ELSE
         YEAR=USERX
      ENDIF
      RETURN
      END
```

```
REAL*4 FUNCTION BENDEL1 (A, E)
C
        Function evaluates the value of the proton-upset cross-section
C
        at energy E using the Bendel 1-parameter inputs
С
С
                         Bendel parameter A
С
         Inputs: A:
                         proton energy (in MeV)
                  E:
С
         Output: SEU cross-section in 10E-12 cm2/bit
С
С
C
C
        Written by:
                       Allan J. Tylka
                       Code 7654
С
                       Naval Research Laboratory
C
                       Washington, DC 20375-5352
C
                       tylka@crs2.nrl.navy.mil
C
C
                       29 March 1996
        Last update:
C
С
С
        IMPLICIT NONE
        REAL*4 Y,A,E
        BENDEL1 = 0.
        Y = (SQRT(18./A)) * (E-A)
        IF (Y.LT.0.) Y=0.
        BENDEL1 = ((24./A)**14.)*((1.-EXP(-.18*SQRT(Y)))**4.)
        IF (BENDEL1.LT.O.) BENDEL1=0.0
        RETURN
        END
```

```
REAL*4 FUNCTION BENDEL2(A,B,E)
С
        Function evaluates the value of the proton-upset cross-section
С
        at energy E using the Bendel 2-parameter inputs
C
C
         Inputs: A, B: Bendel parameters A and B
С
                         proton energy (in MeV)
                  E:
C
         Output: SEU cross-section in 10E-12 cm2
С
C
                       Allan J. Tylka
С
        Written by:
                       Code 7654
C
                       Naval Research Laboratory
C
                       Washington, DC 20375-5352
С
                       tylka@crs2.nrl.navy.mil
С
С
        Last update: 29 March 1996
С
C
        IMPLICIT NONE
        REAL*4 Y, A, B, E
        BENDEL2=0
        Y = (SQRT(18./A)) * (E-A)
        IF (Y.LT.O.) Y=0.
        BENDEL2 = ((B/A) **14.) *((1.-EXP(-.18*SQRT(Y))) **4.)
        IF (BENDEL2.LT.O.) BENDEL2=0.0
        RETURN
        END
```

```
subroutine blccoords(lat, lon, alt, year, imod, bob0, L, yearp, B)
      Inputs:
C
C
          lat, latitude
          lon, east longitude
С
          alt, altitude in km
C
          year, not currently used
C
           imod, = 1 for solar min model, = 2 for solar max model (these are the
C
                  only choices provided for this first version of blccoords)
C
      Outputs:
C
           bob0 returned as Tylka requested (see caveat below where calculated)
С
C
           yearp, year used by allmag (1964 for solar min, 1970 for solar max)
C
C
C**********************
      implicit none
      save
     real*4 lat, lon, alt, year, bob0, L, yearp, b
      real*8 lat8, lon8, alt8, err8, L8, b8
      integer *4 imod, imodold, model
      real*8 constem
      common /gmagmo/ constem
      data err8/ .1/
      data imodold /-10/
      if (imodold .eq. -10) then ! first time
         imodold = imod
        if (imod .eq. 1) then
           yearp = 1964
        elseif (imod .eq.2) then
           yearp = 1970
         else
            stop 'blccords0'
        endif
        call stmag(imod, yearp)
      else
         if (imod. ne. imodold) then
            !imod has changed
           stop 'blccords1'
        endif
      endif
     lat8 = dble(lat)
      lon8 = dble(lon)
     alt8 = dble(alt)
      call invara (imod, yearp, lat8, lon8, alt8, err8, b8, L8)
     b = sngl(b8)
     L = sngl(18)
      compute b over b0 for Tylka....this may not be the bob0 value computed and
C
      used by the trapped proton models because of checks on data base limits.
С
      As an alternative, one can return actual value used from call to
C
      subroutine Trapped protons
C
     bob0 = (b*(L*L*L) / sngl(constem))
```

```
return
     end
     subroutine allmag(model, tm, rkm, st, ct, sph, cph, br, bt, bp, b)
C**********************************
     implicit none
     save
     real*8 ar, aor, b, br, bt, bp, const, cp, cph, ct, dp
     real*8 fn, fm, g, p, par, rkm, sp, sph, st, temp
     real * 4 tm
     integer*4 nmax, m, model, n
     common/magcof/ g(14,14), fn(14), fm(14), const(14,14), nmax
     dimension p(14,14), dp(14,14), sp(14), cp(14)
     data p(1,1), cp(1), dp(1,1), sp(1) / 2*1.,2*0. /
     sp(2) = sph
     cp(2) = cph
     do m = 3, nmax
         sp(m) = sp(2) * cp(m-1) + cp(2) * sp(m-1)
         cp(m) = cp(2) * cp(m-1) - sp(2) * sp(m-1)
     enddo
     aor = 6371.2/rkm
     ar = aor * aor * aor
     p(2,1) = ct
     dp(2,1) = -st
     p(2,2) = st
     dp(2,2) = ct
     br = -(ar+ar)*(g(2,1)*p(2,1)+p(2,2)*(g(2,2)*cp(2)+g(1,2)*sp(2)))
     bt = ar * (g(2,1)*dp(2,1)+dp(2,2)*(g(2,2)*cp(2)+g(1,2)*sp(2)))
     bp = ar * (g(1,2) * cp(2) - g(2,2) * sp(2)) * p(2,2)
     do n = 3, nmax
        ar = aor*ar
         do m = 1, n
            if(m.ne.n) then
               p(n,m) = ct * p(n-1,m) - const(n,m) * p(n-2,m)
               dp(n,m) = ct * dp(n-1,m) - st*p(n-1,m) - const(n,m)*dp(n-2,m)
             else
               p(n,n) = st * p(n-1,n-1)
               dp(n,n) = st * dp(n-1,n-1) + ct * p(n-1,n-1)
             endif
            par = p(n,m) * ar
            if(m.ne.1) then
               temp = g(n,m) * cp(m) + g(m-1,n) * sp(m)
              bp = bp - (g(n,m)*sp(m)-g(m-1,n)*cp(m)) * fm(m) * par
               temp = g(n,m)
            endif
            br = br - temp * fn(n) * par
           bt = bt + temp * dp(n,m) * ar
         enddo
     enddo
     bp = bp/st/100000.
     br = br/100000.
     bt = bt/100000.
```

else

```
IF (abs(bp) .LT. 1.0E18 .AND. abs(br) .LT. 1.0E18 .AND.
        abs(bt) .LT. 1.0E18) THEN
       b = sqrt(br*br + bt*bt + bp*bp)
     ELSE
       b = 1.0E18
     ENDIF
     return
     end
C******************************
     subroutine carmla (B, xi, vl)
C************************
     implicit none
     save
     real*8 B, constem, gg, vl, xi, xx
     compute 1
С
     *************
C
     Equations Containing Constant Mag Moment Will Be Commented
C
C
     Out And Rewritten With New Mag Moment Calculated In Stmag
C
                  3/15/91
     Subroutine
     *****************
C
     common /gmagmo/ constem
     if( xi-1.0d-36 .le. 0.) then
       vl = (constem/B) ** (1./3.)
       return
     endif
     xx = 3.0 * dlog(xi)
     xx = xx + dlog(B/constem)
     if (xx+22. .le. 0.) then
        gg = .333338*xx+.30062102
        go to 7
     endif
     if (xx+3. .le. 0.) then
       gg=(((((((((-8.1537735d-14*xx+8.3232531d-13)*xx+1.0066362d-9)*xx+
    1 8.1048663d-8) *xx+3.2916354d-6) *xx+8.2711096d-5) *xx
    2 +1.3714667d-3)*xx+.015017245)*xx+.43432642)*xx+.62337691
       go to 7
     endif
     if (xx-3. .le. 0.0) then
        gg=((((((((((2.6047023d-10*xx+2.3028767d-9)*xx-2.1997983d-8)*xx-
        5.3977642d-7) *xx-3.3408822d-6) *xx+3.8379917d-5) *xx +
        1.1784234d-3)*xx+1.4492441d-2)*xx+.43352788)*xx+.6228644d0
        go to 7
     endif
     if(xx-11.7 .le. 0.) then
        gg=((((((((((6.3271665d-10*xx-3.958306d-8)*xx+9.9766148d-07)*xx-
        1.2531932d-5) *xx+7.9451313d-5) *xx-3.2077032d-4) *xx +
        2.1680398d-3)*xx+1.2817956d-2)*xx+.43510529)*xx+.6222355d0
        go to 7
     endif
     if (xx-23. .le. 0) then
        gg=(((((2.8212095d-8*xx-3.8049276d-6)*xx+2.170224d-4)*xx -
        6.7310339d-3) *xx+.12038224) *xx-.18461796) *xx+2.0007187
```

```
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```

```
gg=xx-3.0460681
     endif
   7 \text{ vl} = (((1.0 + \text{dexp (gg)}) * \text{constem})/B) * * (1./3.)
     end compute 1
     return
     end
subroutine intega(arc, beg, bend, b, jep, eco, fi)
C************************
     IMPLICIT REAL*8 (A-H, O-Z), INTEGER (I-N)
C
     implicit none
     save
     real*8 a, arc, arg1, asum, b, bb, beg, bend, c, dn, eco, fi
     real*8 t, tb, te, x2, x3
     integer*4 i, kk, jep
     dimension arc(200), beg(200), bend(200), b(200), eco(200)
C
      dimension arc(1), beg(1), bend(1), b(1), eco(1)
     kk = jep
      if (kk .gt. 4) go to 20
      if (kk .eq. 4) kk = kk-1
     a = b(kk-1)/b(2)
     x2 = b(kk)/b(2)
     x3 = b(kk+1)/b(2)
     asum = arc(kk) + arc(kk+1)
      dn = arc(kk) * arc(kk+1) * asum
     bb = (-a*arc(kk+1) * (arc(kk)+asum)+x2*asum**2-x3*arc(kk)**2) /dn
      c = (a*arc(kk+1) - x2 * asum + x3 * arc(kk))/dn
      fi = .157079632d+01 * (1.0-a+bb*bb/(4.0*c)) / dsqrt(dabs(c))
      return
   20 t = dsqrt(1.0d0-bend(2)/b(2))
      fi = (2.0*t-dlog((1.0+t)/(1.0-t)))/eco(2)
      if (b(2)-bend(kk) .gt. 0.) kk=kk+1
      t = dsqrt(dabs(1.0-beg(kk)/b(2)))
      fi = fi-(2.0*t-dlog((1.0+t)/(1.0-t)))/eco(kk)
      kk = kk - 1
   22 do i = 3, kk
         arg1 = 1. - bend(i)/b(2)
         if (arg1 .le. 0.) then
            te = 1.d-5
         else
            te = dsqrt(arg1)
         endif
         arg1 = 1. - beg(i)/b(2)
         if (arg1 .gt. 0.) then
            tb = dsqrt(arg1)
         else
            tb = 1.d-5
         endif
         if (dabs(eco(i))-2.d-5 .le. 0.) then
            fi = fi + ((te+tb)*(arc(i)+arc(i+1)))/4.
            fi = fi+(2.*(te-tb)-dlog((1.+te)*(1.-tb)/((1.-te)*(1.+tb))))
            /eco(i)
     1
         endif
      enddo
```

į.

4

```
30 return
      end
       subroutine invara (model, tm, flat, flong, alt, err, bb, fn)
C**********************************
C **** Note, Error In L Is Typically Less Than 10.*Err*L (Percent)
C **** Flat=Latitude In Degrees , Flong=Longitude In Degrees
C **** Alt=Altitude=Distance From Surface Of Earth In Kilometers
      IMPLICIT REAL*8 (A-H, O-Z), INTEGER (I-N)
      implicit none
      save
      real*4 Tm
     real*8 alt, arc, asum, b, bb, bco, beg, bend, blog, cco, dclt
      real*8 dco, dn, dx, eco, err, fl, flint
     real*8 flat, flong, fn, r1, r2, r3, sa, sc, v, vp, vn
      integer*4 i, j, jep, jup, model
     dimension v(3,3), b(200), arc(200), vn(3), vp(3), beg(200),
     1 bend(200), blog(200), eco(200), r1(3), r2(3), r3(3)
     v(1,2) = alt/6371.2
     v(2,2) = (90.-flat)/57.2957795
     v(3,2) = flong/57.2957795
     arc(1) = 0.
     arc(2) = (1.0+v(1,2)) * sqrt(err) * 0.3
     dclt = 1.5708-0.2007 * dcos(v(3,2) + 1.239)
     if (v(2,2) .gt. dclt) arc(2) = -arc(2)
     call starta(r1, r2, r3, b, arc, v, model, tm)
     do i = 1, 3
        vp(i) = v(i,2)
        vn(i) = v(i,3)
     enddo
     call linesa(r1, r2, r3, b, arc, err, j, vp, vn, model, tm)
     if (j .ge. 200) then
         fl = -1.0
         go to 18
     endif
     iup = i
     do j =1, jup
        arc(j) = dabs(arc(j))
        blog(j) = dlog(b(j))
     enddo
      jep = jup-1
     do j = 2, jep
        asum = arc(j) + arc(j+1)
        dx = blog(j-1) - blog(j)
        dn = asum * arc(j) * arc(j+1)
        bco = ((blog(j-1) - blog(j+1)) * arc(j)**2 -dx *asum**2)/dn
        cco = (dx * arc(j+1) - (blog(j) - blog(j+1)) * arc(j)) / dn
        sa = .75 * arc(j)
        sc = sa + .25 * asum
        dco = blog(j-1) - cco * sa * sc
        eco(j) = bco + cco * (sa + sc)
        beg(j) = dexp(dco+eco(j) * .5 * arc(j))
        bend(j) = dexp(dco+eco(j) * .5 * (asum+arc(j)))
     enddo
     beg(jup) = bend(jep)
     bend(jup) = b(jup)
     eco(jup) = (2.0/arc(jup)) * dlog(bend(jup)/beq(jup))
```

```
call intega(arc, beg, bend, b, jep, eco, flint)
     call carmla (b(2), flint, fl)
  18 bb = b(2)
     fn = fl
     return
     end
            ************
     subroutine linesa(r1, r2, r3, b, arc, err, j, vp, vn, model, tm)
implicit real*8(a-h,o-z),integer(i-n)
     implicit none
     save
     real*4 tm
     real*8 a1, a2, a3, aa, aab, ad, am, ao6, arc, arcj, asum, b
     real*8 bb, bd, bp, br, bt, cc, cd, cop, cot, cre, dd, dn, err
     real*8 pre1, pre2, pre3, qrt, r1, r2, r3
     real*8 ra, rbar, rkm, rt, sip, sit, sna, ssq, vp, vn, x
     integer*4 i, ilp, is, j, model
     dimension b(200), arc(200), r1(3), r2(3), r3(3), vn(3), vp(3), ra(3)
C
     dimension b(1), arc(1) !subroutine arguments
     dimension r1(3), r2(3), r3(3), vn(3), vp(3), ra(3)
     cre = 0.25
     if (err .lt. 0.15625) cre = (err**0.333333333)
     a3 = arc(3)
     aab = dabs(a3)
     sna = a3/aab
     a1 = arc(1)
     a2 = arc(2)
     ao6 = a3*a3/6.0
     j = 3
     ilp = 1
     is = 1
     GO TO 87
   66 \text{ is} = 1
     j = j+1
     ao6 = a3 * a3/6.0
     arcj = a1 + a2 + a3
     ad = (asum+a1)/aa
     bd = asum/bb
     cd = a1/cc
   36 \text{ do i} = 1, 3
        dd = r1(i)/aa-r2(i)/bb+r3(i)/cc
        if (is .eq. 1) then
          rt = r1(i) - (ad*r1(i)-bd*r2(i)+cd*r3(i)-dd*arcj) * arcj
           ra(i) = r1(i)
           r1(i) = r2(i)
           r2(i) = r3(i)
           r3(i) = rt
           vp(i) = vn(i)
        rbar = (r2(i) + r3(i))/2. - dd * ao6
         vn(i) = vp(i) + a3 * rbar
      enddo
   87 if (vn(2) .lt. 0.) vn(2) = -vn(2)
```

```
77 if (vn(2) .le. 3.141592653) go to 78
    vn(2) = 6.283185307 - vn(2)
    go to 77
 78 if (vn(3) .ge. 0.) go to 81
    vn(3) = vn(3) + 6.283185307
    go to 78
 81 if(vn(3) .le. 6.283185307) go to 82
    vn(3) = vn(3) - 6.283185307
    go to 81
82 go to (9, 10), is
  9 \text{ sit} = \text{dabs}(\text{dsin}(\text{vn}(2)))
    prel = vn(1)
    pre2 = pre1 * vn(2)
    pre3 = pre1 * sit * vn(3)
100 \text{ rkm} = \text{vn}(1) * 6371.2
    IF (rkm .lt. 100.0) rkm=100.0
    ssq = sit * sit
    cot = dcos(vn(2))
    sip = dsin(vn(3))
    cop = dcos(vn(3))
    call allmag(model, tm, rkm, sit, cot, sip, cop, br, bt, bp, b(j))
    Added error checking on b(j), 11-24-97.
    IF (b(j) .EQ. 0.0) b(j) = 1.0E-10 !avoid underflows, 11-24-97.
      r3(1) = br/b(j)
      dn = b(j) * vn(1)
      r3(2) = bt/dn
      r3(3) = bp/(dn*sit)
    asum = a3+a2
    aa = asum*a2
    bb = a3*a2
    cc = asum*a3
    is = 2
    go to 36
 10 sit = dabs(dsin(vn(2)))
    b(j) = b(j) * ((pre1/vn(1))**3)
    qrt = 0.5d0 * dabs(r3(1))/(0.1d0+dabs(r3(2)*vn(1)))
    x = (dabs(vn(1)-pre1)+qrt*dabs(vn(1)*vn(2)-pre2)+dabs(vn(1)*sit*
   1 vn(3)-pre3))/(aab*err*dsqrt(1.+qrt*qrt))
    go to (90, 93, 90), ilp
 93 if (x - 3.3) 90, 89, 89
 89 \ a3 = a3 * 0.2 * (8.0+x)/(0.8+x)
    j = j - 1
    ilp = 3
    asum = a2+a1
    aa = asum * a1
    bb = a2 * a1
    cc = asum * a2
    do i = 1, 3
       vn(i) = vp(i)
       r3(i) = r2(i)
       r2(i) = r1(i)
       r1(i) = ra(i)
    enddo
    go to 73
```

```
90 if (j .gt. 200) go to 60
        a1 = a2
        if(b(j)-b(2) .gt. 0) go to 60
           if(b(j)-b(2) .gt. 0 .OR. b(j) .EQ. 0.0) go to 60 !changed 11-24-97.
  CF?
        ilp = 2
        a2 = a3
        a3 = a3 * .2 * (8.+x)/(.8+x)
        am = (2.-r3(2) * vn(1)) * vn(1) * cre
        if (dabs(a3) - am .gt. 0.) a3 = sna * am
        if(sna * r3(1)+.5 .gt. 0.) go to 73
        am = -.5 * sna * vn(1)/r3(1)
        if (dabs(a3) - am .gt. 0) a3 = sna * am
    73 \text{ arc}(j+1) = a3
        aab = dabs(a3)
        go to 66
     60 return
        end
  subroutine starta(r1, r2, r3, b, arc, v, model, tm)
  implicit real*8(a-h,o-z),integer(i-n)
        implicit none
w.
        save
        real*8 aer, arc, b, bp, br, bt, cop, cot,
        real*8 oer, sip, sit, ssq, r1, r2, r3, rkm, v
        real*4 tm
Li
        integer*4 i, is, model
dimension b(1), arc(1), v(3, 3), r1(3), r2(3), r3(3)
  С
Ξ
        dimension b(1), arc(1), v(3, 3), r1(1), r2(1), r3(1)
                                                          ! arguments
i.
P.
        sit = dabs(dsin(v(2,2)))
L
        aer = v(1,2)
L.
        ssq = sit * sit
        oer = (6356.912 + ssq * (21.3677 + .108 * ssq))/6371.2
        v(1,2) = aer + oer
     10 if (v(3,2) .ge. 0. ) go to 12
        v(3,2) = v(3,2) + 6.283185307
        go to 10
     12 \text{ rkm} = v(1,2) * 6371.2
        if (model.eq.6) rkm = rkm+14.288-ssq * (21.3677+.108 * ssq) !CHECK ORIG MODEL NO; 'S
        cot = dcos(v(2,2))
        sip = dsin(v(3,2))
        cop = dcos(v(3,2))
        call allmag (model, tm, rkm, sit, cot, sip, cop, br, bt, bp, b(2))
        r2(1) = br/b(2)
        dn = b(2) * v(1,2)
        r2(2) = bt/dn
        r2(3) = bp/(dn * sit)
        is = 0
      1 \text{ do } i = 1, 3
           v(i,1) = v(i,2) - arc(2) * r2(i)
        sit = dabs(dsin(v(2,1)))
      3 \text{ rkm} = v(1,1) * 6371.2
```

```
ssq = sit * sit
        if(model.eq.6) rkm = rkm+14.288-ssq * (21.3677+.108 * ssq)
        cot = dcos(v(2,1))
        sip = dsin(v(3,1))
        cop = dcos(v(3,1))
        call allmag (model, tm, rkm, sit, cot, sip, cop, br, bt, bp, b(1))
        if (b(1)-b(2) .ge. 0.) go to 5
        arc(2) = -arc(2)
        go to 1
      5 r1(1) = br/b(1)
        arc(3) = arc(2)
        dn = b(1) * v(1,1)
        r1(2) = bt/dn
        r1(3) = bp/(dn * sit)
        do i=1, 3
           v(i,1) = v(i,2) - arc(2) * (r1(i) + r2(i))/2.
        sit = dabs(dsin(v(2,1)))
        is = is+1
        go to (3, 7), is
  C
        if (is .eq. 1) go to 3
      7 \text{ do i} = 1, 3
        do i = 1, 3
           v(i,3) = v(i,2) + arc(3) * ((1.5) * r2(i) - .5 * r1(i))
enddo
        return
        end
        subroutine stmag(model, tm)
    *********************
  C
        Constant Mag Moment Replaced With Calculated Mag Moment,
         Using Geomagnetic Field Expansion Coefficients - 3/15/91
  C
52
k
                        choice of 2 models - see below
       Inputs model
  C
geocentric distance in kilometers
               rkm
  C
time in years for desired field
  C
               tm
į.
                        sin + cos of geocentric colatitude
  C
               st,ct
1
  C
               sph,cph sin + cos of east longitude
  C
      Outputs br, bt, bp geocentric field components in gauss
                        field magnitude in gauss
  C
        IMPLICIT REAL*8(A-H,O-Z)
  С
        implicit none
        save
        real*8 const, constem, em, f1, f2, f3, fm, fn, g
        real*8 rad, t, t0
        real*4 tm, tmold
        integer*4 jj, k, l, m, modold, n, nmax, nmx, model
        common /gmagmo/ constem
        integer*4 g1(13,13),gt1(13,13),gtt1(13,13), g2(13,13), gt2(13,13),
       1 gtt2(13,13), lg(13,13,2), lgt(13,13,2), lgtt(13,13,2)
        real*4 gg(13,13,2), ggt(13,13,2), ggtt(13,13,2), shmit(13,13)
        equivalence (g1,gg,lg), (gt1,ggt,lgt), (gtt1,ggtt,lgtt),
              (g2, lg(1,1,2)), (gt2, lgt(1,1,2)), (gtt2, lgtt(1,1,2))
```

```
character*32 label(2)
      dimension t0(2), nmx(2)
      common/magcof/ g(14,14), fn(14), fm(14), const(14,14), nmax
      data label(1)/'IGRF 1965.0 80-TERM 10/68
      data label(2)/'HURWITZ US C+GS 168-TERM 1970 '/
      data t0/1965.d+00,1970.d+00/
      data nmx/9,13/
C ***** G1,Gt1 Igrf 1965.0
                                          80-Term
                                                        10/68 Epoch 1965
      data g1 / 1, -30339,-1654,1297,958,-223,47,71,10,4*0,5758,-2123,
     A 2994,-2036,805,357,60,-54,9,4*0,-2006,130,1567,1289,492,246,4,0,
     B -3,4*0,-403,242,-176,843,-392,-26,-229,12,-12,4*0,149,-280,8,-265
     C ,256,-161,3,-25,-4,4*0,16,125,-123,-107,77,-51,-4,-9,7,4*0,-14,
     D 106,68,-32,-10,-13,-112,13,-5,4*0,-57,-27,-8,9,23,-19,-17,-2,12,
     E 4*0,3,-13,5,-17,4,22,-3,-16,6,56*0/
      data gt1 / 10, 153,-244,2,-7,19,-1,-5,1,4*0,-23,87,3,-108,2,11,-3,
     F = 3,4,4*0,-118,-167,-16,7,-30,29,11,-7,6,4*0,42,7,-77,-38,-1,6,19,
     G -5,5*0,-1,16,29,-42,-21,0,-4,3,5*0,23,17,-24,8,-3,13,-4,0,-1,4*0,
     H-9, -4, 20, -11, 1, 9, -2, -2, 3, 4*0, -11, 3, 4, 2, 4, 2, 3, -6, -3, 4*0, 1, -2, -3, -2,
     I -3, -4, -3, -3, -5, 56*0/
      data gtt1 /1,168*0/
  ***** G2,GT2 Hurwitz Us Coast + Geodetic S. 168-Term
                                                               Epoch 1970.
      data g2/10,-302059,-17917,12899,9475,-2145,460,734,121,107,-39,16,
     A -4,57446,-20664,29971,-20708,8009,3595,651,-546,77,57,-26,-31,30,
     B -20582,430,16086,12760,4579,2490,95,46,-32,23,7,-36,5,-3699,2456,
      \texttt{C-1880,8334,-3960,-290,-2188,175,-124,-110,-19,37,-3,1617,-2758,} \\
     D 185,-2788,2436,-1669,20,-210,-44,131,-15,-3,-13,157,1420,-1310,
     E = -911, 808, -582, -22, -32, 45, 33, 74, -6, 4, -171, 1146, 625, -323, -78, 38,
     F -1125,143,34,2,46,-8,-14,-666,-265,-34,81,209,-240,-186,41,125,
     G 15,6,1,-12,121,-160,22,-176,46,189,-46,-187,94,9,-8,2,-12,-174,
     H 163,14,-27,-32,80,137,-4,-14,-4,22,-24,-1,27,19,0,35,-45,22,-31,
     I 56,-1,-63,14,4,10,-2,26,-26,-9,21,-1,18,-14,-28,-17,-14,6,-4,-3,
     J 4,9,-1,-10,26,-32,13,-6,-19,7,19,12/
      data gt2/10,231,-244,-19,-7,12,-7,0,3,4*0,-46,112,-1,-90,-6,7,6,
     K = 3, 3, 4*0, -104, -166, 40, -20, -36, 12, 14, 3, 4, 4*0, 72, 21, -52, -54, -11, 0,
     L 17,6,1,4*0,22,-5,14,-24,-23,-15,6,3,-1,4*0,1,25,-14,9,1,11,-3,2,
     M = 3,4*0,-5,11,2,-3,7,22,-5,1,9,4*0,-17,-3,7,1,-2,-3,-2,-1,-2,4*0,
     N = 2, -6, -3, -4, 1, -2, -2, -1, 6, 56*0/
      data gtt2 /1,168*0/
      data shmit(1,1) / 0.0 /, tmold / -100./
      parameter (rad = 57.29577636718750)
C ***** Begin Program
      if(model .lt. 1 .or. model .gt. 2) stop 'stmag1'
      if(shmit(1,1).eq.-1.) go to 8 ! already initialized
      do n = 1, 14
         fn(n) = n
         do m = 1, 14
            fm(m) = m-1
            const(n,m) = float((n-2)**2-(m-1)**2) / ((2*n-3)*(2*n-5))
```

```
and the same time that the many time as the same time time to the same time to the same time.
```

```
enddo
    enddo
        Initialize * Once Only, First Time Subroutine Is Called
****
    shmit(1,1) = -1.
    do n = 2, 13
       shmit(n,1) = (2 * n-3) * shmit(n-1,1) / (n-1)
       jj=2
       do m = 2, n
          shmit(n,m) = shmit(n,m-1) *sqrt(float((n-m+1) * jj)/(n+m-2))
          shmit(m-1,n) = shmit(n,m)
          jj = 1
       enddo
    enddo
   do k = 1, 2
      f1 = lg(1,1,k)
      f2 = lgt(1,1,k)
      f3 = lgtt(1,1,k)
      nmax = nmx(k)
      1 = 0
      do n = 1, nmax
         do m = 1, nmax
            gg(n,m,k) = lg(n,m,k) * shmit(n,m)/f1
            ggt(n,m,k) = lgt(n,m,k) * shmit(n,m)/f2
            ggtt(n,m,k) = lgtt(n,m,k) * shmit(n,m)/f3
         enddo !m
      enddo !n
   enddo !k
 8 if (model.eq.modold) return
***** NOTE WRITE STATEMENT - NEW MODEL OR NEW TIME
   type 9,
            model, label ( model) ,tm
 9 format(' model used is number',i2,2x,a32,' for tm =',f9.3/)
   modold = model
   tmold = tm
   nmax = nmx(model)
   t = tm-t0 \pmod{1}
   do n = 1, nmax
      do m = 1, nmax
        g(n,m) = gg(n,m,model) + t * (ggt(n,m,model) + ggtt(n,m,model) *t)
      enddo
   enddo
   em = sqrt(g(1,2)**2 + g(2,2)**2 + g(2,1)**2)
   constem = em/100000.0
   return
   end
```

```
SUBROUTINE CALC_SEU_RATE(NBITS,SEU_RATE,DAY_RATE,
                                PERSECOND, PERDAY)
     &
       IMPLICIT NONE
       REAL*4 NBITS, SEU_RATE, DAY_RATE, PERSECOND, PERDAY
C
C
       INPUTS:
C
                 NBITS
                         = number of bits per device
                 SEU_RATE = calculated SEU rate (in upsets/bit/second)
С
C
       OUTPUTS:
                 PERSECOND = upsets/device/second
C
                         = upsets/device/day
C
                 PERDAY
                 DAY_RATE = upsets/bit/day
C
       PERSECOND=NBITS*SEU_RATE
       PERDAY=PERSECOND*24.0*3600.
       DAY RATE=SEU RATE*24.0*3600.
```

RETURN

END

```
SUBROUTINE CAPITALIZE_STRING(STRING,ILONG)
C
C
       Re-writes an input character string STRING of length ILONG
C
       into all capitals.
С
       IMPLICIT NONE
       CHARACTER STRING
       INTEGER*4 ILONG, I
       IF (ILONG.GE.1) THEN
           DO 100 I=1, ILONG
           IF (ICHAR(string(i:i)) .GE. 97) THEN
           string(i:i) = CHAR(ICHAR(string(i:i)) - 32)
           ENDIF
           CONTINUE
 100
       ENDIF
       RETURN
       END
```

```
SUBROUTINE CHECK CREME96 VERSION(FILENAME, IVER)
С
      Examines first line of input file, to get version number
C
C
      IMPLICIT NONE
      CHARACTER*80 FILENAME, ILINE
      CHARACTER*3 VERSIONLABEL
      INTEGER*4 NCHAR, IVER, STAT, CREME96 OPEN
      INTEGER*4 FILENO
      DATA FILENO/4/
С
     Modified 7/29/96: Version 1.01
      OPEN (UNIT=FILENO, FILE='USER:'//FILENAME, STATUS='OLD',
С
            READONLY, SHARED)
      stat = creme96_open(filename,'user',fileno,'old')
      READ(FILENO, 1) ILINE
 1
      FORMAT (A80)
      NCHAR=3
      VERSIONLABEL=ILINE (76:78)
      IF (VERSIONLABEL.EQ.' ') THEN
          IVER=0
      ELSE
          read(versionlabel,'(i3)') iver
      ENDIF
      CLOSE (FILENO)
      RETURN
      END
```

```
SUBROUTINE CHECK FILE (IFILETYPE, FILENAME, IACCEPT)
   C
   C
           Subroutine for checking existence and acceptability of specified
   C
           input file.
   C
           IMPLICIT NONE
           INTEGER*4 MHMAX, NLINES, MAXFILETYPE
           PARAMETER (MAXFILETYPE=8)
           CHARACTER*80 ILINES, TEMPLINE
           PARAMETER (MHMAX=20)
           DIMENSION ILINES (MHMAX)
           INTEGER*4 IFILETYPE, JFILETYPE, J, IACCEPT, IANSWER, NHMAX, IHMAX
           DIMENSION IHMAX (MAXFILETYPE)
           DATA IHMAX/3,3,10,15,20,20,4,1/
           CHARACTER*80 FILENAME
           LOGICAL IEXIST, FILE_CHECKS, CREME96_INQUIRE
           INTEGER*4 IERR
           DATA IERR/0/
   C
   C
           IACCEPT=0
           CALL GET CHECK CONTROL (FILE CHECKS)
           IF (.not. FILE CHECKS) RETURN
₫ C
           First, see if specified input file exists:
           INQUIRE(FILE='USER:'//FILENAME,EXIST=IEXIST)
T C
           iexist = creme96 inquire(filename,'user')
          IF (.NOT. IEXIST) THEN
T.
               WRITE(6,999)
     999
               FORMAT(1X,' This file was not found in USER area.',
               ' Please try again.')
CALL SHOW DIRECTORY (IFILETYPE)
121
               IACCEPT=-1
lesk.
               RETURN
75
           ENDIF
L.
Ēz.
           IF (IFILETYPE.EQ.0) RETURN
ر
ا
          Now see if file has correct type:
           CALL CHECK FILE TYPE (FILENAME, JFILETYPE)
           IF (JFILETYPE.GT.0) THEN
               IF (IFILETYPE.EQ.1) THEN
                    IF (JFILETYPE.NE.1) THEN
                       WRITE(6,9001)
    9001
                       FORMAT(1x,' This file does not contain a trapped',
                       ' proton flux. Please try again.')
                       CALL SHOW DIRECTORY (IFILETYPE)
                       IACCEPT=-1
                    ENDIF
               ELSEIF (IFILETYPE.EQ.2) THEN
                    IF (JFILETYPE.NE.2) THEN
                       WRITE(6,9002)
                   {\tt FORMAT}\,({\tt lx},{\tt '}{\tt This}\ {\tt file}\ {\tt does}\ {\tt not}\ {\tt contain}\ {\tt a}\ {\tt geomagnetic'},
    9002
                    ' transmission function.',/,1x,' Please try again:')
                       CALL SHOW DIRECTORY (IFILETYPE)
                       IACCEPT=-1
                    ENDIF
               ELSEIF (IFILETYPE.EQ.3 .or. IFILETYPE.EQ.4) THEN
                     IF (JFILETYPE.NE.3 .and. JFILETYPE.NE.4
```

```
WRITE (6,9003)
         9003
                                                  FORMAT(1x,' This file does not contain particle',
                                                  ' fluxes. Please try again.')
                   &
                                                  CALL SHOW DIRECTORY (IFILETYPE)
                                                  IACCEPT=-1
                                               ENDIF
                                 ELSEIF (IFILETYPE.EQ.5) THEN
                                             IF (JFILETYPE.NE.5) THEN
                                                 WRITE(6,9005)
         9005
                                                 FORMAT(1x,' This file does not contain an integral',
                                                  ' LET spectrum. Please try again.')
                                                  CALL SHOW DIRECTORY (IFILETYPE)
                                                  IACCEPT=-1
                                             ENDIF
                                ELSEIF (IFILETYPE.EQ.6) THEN
                                             IF (JFILETYPE.NE.6) THEN
                                                 WRITE(6,9006)
        9006
                                                 FORMAT(1x,' This file does not contain a differential',
                                                 ' LET spectrum. Please try again.')
                  &
                                                 CALL SHOW DIRECTORY (IFILETYPE)
                                                  IACCEPT=-1
                                            ENDIF
                              ELSEIF (IFILETYPE.EQ.7) THEN
                                            IF (JFILETYPE.NE.7) THEN
A CONTROL OF THE PARTY OF THE P
                                                 WRITE(6,9007)
        9007
                                                 FORMAT(1x,' This file does not contain a shielding',
                                                 ' distribution prepared by the CREME96 software.',
                  &
                                             /,' Please try again.')
                                                 CALL SHOW DIRECTORY (IFILETYPE)
Į.
                                                 IACCEPT=-1
ENDIF
ELSEIF (IFILETYPE.EQ.8) THEN
E
                                            IF (JFILETYPE.NE.O .and. JFILETYPE.NE.8) THEN
WRITE (6, 9008)
9008
                                                 FORMAT(1x,' This file does not contain a',
W
                                                 ' cross-section table.',
                  æ
ĿĿ
                                             /,' Please try again.')
CALL SHOW DIRECTORY (IFILETYPE)
٠..[
                                                 IACCEPT=-1
                                            ENDIF
                              ENDIF
                       IF (IACCEPT.LE.-1) RETURN
                      ENDIF
                      IF (IFILETYPE.GT.O .and. IFILETYPE.LE.MAXFILETYPE) THEN
                      NHMAX=IHMAX (IFILETYPE)
                      CALL UNLOAD HEADERS (FILENAME, NHMAX, ILINES, NLINES)
                      DO 100 J=1, NLINES
                              TEMPLINE=ILINES (J)
                              IF (TEMPLINE(2:2).EQ.'%') WRITE(6,997) TEMPLINE
        997
                              FORMAT (A80)
                      CONTINUE
        100
                      ENDIF
        101
                      CONTINUE
                      CALL RETRY INPUT (IERR)
                      WRITE (6, 996)
        996
                      FORMAT(1x,' Is this the input file you want here? 0=no,1=yes')
```

READ(*, *, ERR=101, IOSTAT=IERR) IANSWER

.and. JFILETYPE.NE.1) THEN

æ

```
PROGRAM CHECK FILE DRIVER
 C
 C
        Stand-alone version of the CHECK_FILE routine, primarily written
 C
        for interface to the WWW version of CREME96.
 C
С
        In response to questions, USER supplies the following information
C
          IFILETYPE: indicates desired type of file (1=.trp, etc.)
C
          FILENAME: filename (without directory name appended).
C
C
       Outputs from the program which are to be subsequently displayed by
C
       the WWW interface are prefaced with preface "**")
C
       IMPLICIT NONE
       INTEGER*4 IFILETYPE, JFILETYPE
       CHARACTER*80 FILENAME
       LOGICAL IEXIST, CREME96 INQUIRE
       INTEGER*4 IERR, K
       CHARACTER*30 DESCRIP
       DIMENSION DESCRIP(11)
       DATA DESCRIP/' a trapped proton flux',
                   ' a geomag transmission fcn',
     &
                    ' external particle fluxes',
     &
                    ' internal particle fluxes',
                    ' an integral LET spectrum',
     &
                   ' a differential LET spectrum',
                    ' a shielding distribution', '
     &
                   ' a cross-section table',
     &
                   ' a PUP-SEU report',
     &
                    ' a HUP-SEU report',
                   ' a dose report'/
      INTEGER*4 NHMAX, LINEMAX
      PARAMETER (NHMAX=30)
       CHARACTER*80 HEADER LINE
       DIMENSION HEADER_LINE(NHMAX)
       CHARACTER*80 ILINE, ILINEOUT
                 ------
C
C
       Get inputs from user:
 101 CONTINUE
       WRITE(6,1000)
 1000 FORMAT(' Specified desired filetype: ',
            /,' 1=.TR*; 2=.GT*; 3=.FLX; 4=.TFX; 5=.LET; 6=.DLT;',
                ' 7=.SHD; 8=.XSD;',
     æ
                9=.PUP; 10=.HUP, 11=.DSE')
       READ(*,*,ERR=101,IOSTAT=IERR) IFILETYPE
 102
       CONTINUE
       IF (IFILETYPE.LT.1 .or. IFILETYPE.GT.11) THEN
          WRITE(6,1005) IFILETYPE
          FORMAT(1x,'ERROR: IFILETYPE = ',18,' not defined. Try again.')
 1005
          STOP
       ENDIF
       WRITE (6, 1100)
 1100 FORMAT (' Enter FILENAME: ')
```

C-----C

C

Now begin analysis of file:

READ(*,105,ERR=102,IOSTAT=IERR) FILENAME

105 FORMAT (A80)

```
First, see if specified input file exists:
  C
          INOUIRE(FILE='USER:'//FILENAME,EXIST=IEXIST)
  С
         iexist = creme96_inquire(filename,'user')
         IF (.NOT. IEXIST) THEN
             WRITE(6,9101) FILENAME(1:78)
             FORMAT (1x, '**', A78,
   9101
                   /,1x,'**not found in USER area. Try again.')
       ક
             STOP
         ELSE
             WRITE(6,9102) FILENAME(1:78)
             FORMAT (1x, '**', A78)
   9102
         ENDIF
  С
         Now see if filetype matches requested type.
          IF (IFILETYPE.EQ.8) THEN
              WRITE(6,9105)
              FORMAT(1x,'**According to the extension (.xsd), this file',
   9105
                   /,1x,'**contains a user-supplied cross-section table.',
                   /,lx,'**Here are the first 2 lines of the file: ')
         ELSEIF (IFILETYPE.NE.8) THEN
              CALL CHECK_FILE_TYPE (FILENAME, JFILETYPE)
IF (IFILETYPE.EQ.JFILETYPE) THEN
                  WRITE(6,9200) DESCRIP(IFILETYPE)
                  FORMAT(1x,'**This file contains',A30,'.')
   9200
              ELSEIF (IFILETYPE.NE.JFILETYPE) THEN
                  WRITE(6,9201) DESCRIP(IFILETYPE)
              FORMAT(1x,'**This file does NOT contain', A30,'. Try again:')
   9201
į.
          ENDIF
ENDIF
Now get header information:
LINEMAX=0
CALL UNLOAD HEADERS (FILENAME, NHMAX, HEADER_LINE, LINEMAX)
          IF (LINEMAX.LE.O) THEN
                 WRITE(6,9301)
                 FORMAT(1x, '**No header information stored in file.')
    9301
          ELSE
                 IF (IFILETYPE.NE.8) WRITE(6,9302)
                 FORMAT(1x,'**Header information stored in this file:')
    9302
                 DO 9400 K=1, LINEMAX
                    ILINE=HEADER LINE(K)
                    ILINEOUT='**'//ILINE(3:80)
                    WRITE(6,9399) ILINEOUT(1:80)
                    FORMAT(1x, A80)
    9399
                 CONTINUE
    9400
          ENDIF
          STOP
          END
```

С

```
SUBROUTINE CHECK_FILE(IFILETYPE, FILENAME, IACCEPT)
С
C
       Subroutine for checking existence and acceptability of specified
C
       input file.
C
       IMPLICIT NONE
       INTEGER*4 MHMAX, NLINES, MAXFILETYPE
       PARAMETER (MAXFILETYPE=8)
       CHARACTER*80 ILINES, TEMPLINE
       PARAMETER (MHMAX=20)
       DIMENSION ILINES (MHMAX)
       INTEGER*4 IFILETYPE, JFILETYPE, J, IACCEPT, IANSWER, NHMAX, IHMAX
       DIMENSION IHMAX (MAXFILETYPE)
       DATA IHMAX/2,3,10,15,20,20,4,1/
       CHARACTER*80 FILENAME
       LOGICAL IEXIST, NO CHECKS, CREME 96 INQUIRE
       DATA NO_CHECKS/.FALSE./
       INTEGER*4 IERR
       DATA IERR/0/
C
```

IACCEPT=0
IF (NO_CHECKS) RETURN

RETURN END

```
SUBROUTINE CHECK_FILE_TYPE(FILENAME, IFILETYPE)
C
      Examines first line of input file, to check file-type code.
C
C
      IMPLICIT NONE
      CHARACTER*80 FILENAME
      INTEGER*4 IVER, STAT, CREME96_OPEN
      INTEGER*4 IFILETYPE, FILENO
      DATA FILENO/4/
      CALL CHECK_CREME96_VERSION(FILENAME, IVER)
      IFILETYPE=0
      IF (IVER.EQ.0) RETURN
      OPEN(UNIT=FILENO, FILE='USER:'//FILENAME, STATUS='OLD',
С
            READONLY, SHARED)
      stat = creme96_open(filename,'user',fileno,'old')
      READ(FILENO, 1) IFILETYPE
      FORMAT (78x, I2)
      CLOSE (FILENO)
      RETURN
      END
```

```
SUBROUTINE CHECK_HEADER_LENGTH(FILENAME, NHEADER)
C
C
      Examines first line of input file, to get header length.
C
      IMPLICIT NONE
      CHARACTER*80 FILENAME, ILINE
      INTEGER*4 IVER, NHEADER, STAT, CREME96 OPEN
      INTEGER*4 FILENO
      DATA FILENO/4/
      CALL CHECK_CREME96_VERSION(FILENAME, IVER)
      IF (IVER.EQ.0.or.IVER.EQ.101) NHEADER=2
      IF (IVER.GE.102) THEN
       OPEN (UNIT=FILENO, FILE='USER:'//FILENAME, STATUS='OLD',
C
            READONLY, SHARED)
      stat = creme96 open(filename, 'user', fileno, 'old')
      READ(FILENO, *) NHEADER
      ENDIF
      CLOSE (FILENO)
      RETURN
      END
```

```
SUBROUTINE CHECK_NAME_CONFLICT(INFILE,OUTFILE,IACCEPT)
Ċ
       Makes sure that input and output names are not identical.
С
С
       IMPLICIT NONE
       CHARACTER*80 INFILE, OUTFILE, TEMPIN, TEMPOUT
       INTEGER*4 IACCEPT, ILONG
       LOGICAL FILE CHECK
       IACCEPT=0
       CALL GET_CHECK_CONTROL (FILE CHECK)
       IF (.not. FILE_CHECK) RETURN
C
       Dispose of version numbers
       ILONG=INDEX(INFILE,';')
       TEMPIN=INFILE
       IF (ILONG.NE.0) TEMPIN=INFILE(1:ILONG-1)
       TEMPOUT=OUTFILE
       ILONG=INDEX(OUTFILE,';')
       IF (ILONG.NE.0) TEMPOUT=OUTFILE(1:ILONG-1)
C
       Convert to upper case:
       ILONG=LEN (TEMPIN)
       CALL CAPITALIZE_STRING (TEMPIN, ILONG)
       ILONG=LEN (TEMPOUT)
       CALL CAPITALIZE_STRING (TEMPOUT, ILONG)
      Now see if identical:
       IF (TEMPIN.EQ.TEMPOUT) THEN
           IACCEPT=-1
           WRITE(6,666) TEMPIN, TEMPOUT
           FORMAT(1x,' INFILE = ',A69,/,1x,' OUTFILE = ',A69,
666
                /,1x,' ERROR: Input and Output filenames are identical.',
    &
                /,1x,' Please enter another output filename.')
      ENDIF
      RETURN
      END
```

```
SUBROUTINE CHECK OUTPUT FILE (FILENAME, IACCEPT)
   С
           Subroutine for checking existence and specified output file.
   С
   C
           IMPLICIT NONE
           CHARACTER*80 FILENAME
           INTEGER*4 IACCEPT, IREPEAT
           LOGICAL IEXIST, FILE CHECK, CREME96_INQUIRE
   C
           IACCEPT=0
           CALL GET CHECK CONTROL (FILE_CHECK)
           IF (.not. FILE CHECK) RETURN
           IEXIST=.FALSE.
          See if specified output file already exists:
   С
           INOUIRE(FILE='USER:'//FILENAME,EXIST=IEXIST)
   C
           iexist = creme96_inquire(filename,'user')
           IF (IEXIST) THEN
               WRITE(6,999)
     999
               FORMAT(1X,' A file with this name',
                          ' already exists in your USER area.',
                     /,lx,' Do you wish to create a new file with',
         æ
                          ' the same name? (0=no,1=yes)')
The time and the limb time that the
               READ(*,*,ERR=101) IREPEAT
               IF (IREPEAT.NE.1) THEN
                 IACCEPT=-1
                 CONTINUE
    101
                 WRITE(6,995)
                 FORMAT(1x,' Please give another name: ')
    995
               ELSE
Ė
                 IACCEPT=0
The three many in
                 WRITE(6,996)
                 FORMAT(1x,' A new file with this same name will',
    996
                             ' be created.')
         æ
               ENDIF
           ENDIF
           RETURN
           END
```

```
&
                                               IPARAM, PARAMS, XSECT FILE,
          &
                                               XM, YM, ZM)
    C
    C
             Routine for extracting lateral RPP dimension from cross-section
    C
             when the input XM, YM values are zero:
    C
    C
             Inputs:
    C
    C
                      XM, YM, ZM = bit dimensions (in microns)
    C
                                   = 1,2,4, indicating cross-section model
    С
                                     1 = Bendel 1-parameter
    C
                                     2 = Bendel 2-parameter
    С
                                     4 = Weibull
    C
                                     5 = Critical charge (pc)
    C
                                     0 = table
    С
                      PARAMS (4)
                                       = array containing cross-section parameters
    C
                      XSECT_FILE = file containing cross-section table.
    C
    С
            Outputs: XM, YM, ZM = revised RPP dimension
    C
    C
    С
            Written by:
                            Allan J. Tylka
    C
                            Code 7654
    С
                            Naval Research Laboratory
C
                            Washington, DC 20375-5352
    C
                            tylka@crs2.nrl.navy.mil
    C
    C-
            IMPLICIT NONE
INTEGER*4 IPARAM, NDUM, NTRY
22
            REAL*4 XMO, YMO, ZMO, PARAMS, XM, YM, ZM, LETMAX, XSMAX, DELTA_XS
1
            CHARACTER*80 XSECT FILE
DIMENSION PARAMS(4), LETMAX(2), XSMAX(2)
IJ
Frei.
            0MX=MX
            YM=YM0
            ZM = ZM0
            IF (XM.GT.1.0E-6 .and. YM.GT.1.0E-6) RETURN
    C
    C
            User has specified XM,YM=0. Must extract value from the
    С
            cross-section inputs:
    C
            IF (IPARAM.NE.5) THEN
            NDUM=2
            NTRY=1
            LETMAX (1) = 1.0E + 5
            LETMAX (2) = 1.0E+6
     10
            CONTINUE
            CALL EVALUATE_SEU_CROSS_SECTION(LETMAX, NDUM, IPARAM, PARAMS,
         &
                                              XSECT FILE, XSMAX)
            DELTA_XS=ABS (XSMAX (1) -XSMAX (2)) /XSMAX (2)
               IF (DELTA XS.GT.0.01) THEN
                  WRITE(6,9999)
                  NTRY=NTRY+1
```

LETMAX (1) = LETMAX (1) *10.0

SUBROUTINE CHECK_RPP_DIMENSIONS(XMO,YMO,ZMO,

```
LETMAX (2) = LETMAX (2) *10.0
             IF (NTRY.LE.10) GOTO 10
          ENDIF
       ELSEIF (IPARAM.EQ.5) THEN
           XSMAX(2)=PARAMS(2)
       ENDIF
       XM=SQRT(XSMAX(2))
       MX=MY
       IF (XM.LT.1.0E-6 .and. YM.LT.1.0E-6) THEN
           WRITE(6,9998) XMO, YMO, XSMAX(2)
           FORMAT (' Error in HUP inputs: ',
9998
                /,' Input lateral RPP dimensions = ',2E13.6,
    &
                /,' Input limiting cross-section = ',E13.6,
    &
                /,' SEU RATE = 0.0 !!!')
       ENDIF
          FORMAT(' ERROR in CHECK_RPP_DIMENSIONS: Plateau not found.')
9999
       RETURN
       END
```

```
UPATH, FRACSHLD,
                                                XMEAN, XRMS, TOTAL, ERRFLAG)
          IMPLICIT NONE
          INTEGER*4 MAXSHIELD
         PARAMETER (MAXSHIELD=500)
         INTEGER*4 NSHIELD, K, INDX (MAXSHIELD)
         REAL*4 UPATHO, FRACSHLDO, UPATH, FRACSHLD
         REAL*4 XMEAN, XRMS, TOTAL
         INTEGER*4 ERRFLAG
         DIMENSION UPATHO(1), FRACSHLDO(1), UPATH(1), FRACSHLD(1)
         IF (NSHIELD.GT.MAXSHIELD) THEN
             WRITE(6,995) NSHIELD, MAXSHIELD
             FORMAT ('@ 07001 ABNORMAL TERMINATION: ',
    995
                   /,1x,' ERROR in CHECK SHIELD DISTRIBUTION: ',
                   /,1x,' TOO MANY BINS: ',18,' > ',18,' max.',
        æ
                   /,1x,' STOP.')
             STOP
         ENDIF
         First, check normalization
         TOTAL=0
         DO 100 K=1, NSHIELD
         TOTAL=TOTAL+FRACSHLD0(K)
    100 CONTINUE
         WRITE(6,999) NSHIELD
    999 FORMAT ('No. Shielding Bins = ', I4)
         WRITE(6,998) TOTAL
    998 FORMAT(' Sum of shielding fractions = ',E13.6)
ERRFLAG=0.0
Œ
         IF (ABS (TOTAL-1.0).GT.0.0001) THEN
E z
               ERRFLAG=1
T.
               WRITE (6,997)
               FORMAT(' Shielding distribution will be re-nomalized',
    997
.
Paris
                       ' to unit integral')
L.
         ENDIF
         XMEAN=0.0
         XRMS=0.0
         DO 200 K=1, NSHIELD
            Renormalize shielding fraction to unit integral:
   C
            FRACSHLDO(K)=FRACSHLDO(K)/TOTAL
   C
            Calculate mean
            XMEAN=XMEAN+UPATHO(K)*FRACSHLDO(K)
   C
            Calculate mean square:
            XRMS= XRMS+FRACSHLD0(K)*UPATH0(K)**2
    200 CONTINUE
         XRMS=XRMS-XMEAN*XMEAN
         IF (XRMS.LT.0.0) XRMS=0.0
         XRMS=SORT (XRMS)
         WRITE(6,250) XMEAN, XRMS
    250 FORMAT(1x,' Mean shielding thickness = ',E13.6,
             /,1x,' RMS deviation
                                              = ',E13.6)
   С
         Now re-order according to increasing shielding thickness. This
         ordering makes the transport code more efficient.
```

SUBROUTINE CHECK_SHIELD_DISTRIBUTION(NSHIELD, UPATH0, FRACSHLD0,

CALL INDEXX (NSHIELD, MAXSHIELD, UPATHO, INDX)

DO 500 K=1, NSHIELD

UPATH(K)

=UPATHO(INDX(K))

FRACSHLD(K) = FRACSHLD0(INDX(K))

500 CONTINUE

RETURN

END

```
SUBROUTINE COPY HEADERS (INFILE, NHEADER, OUTUNIT)
C
C
      Reads NHEADER lines of header information from file INFILE
С
      and copies to unit OUTUNIT (which has previously been opened
С
      by the calling routine).
      CHARACTER*80 INFILE, ILINE
      INTEGER*4 NHEADER, INUNIT, OUTUNIT, IVER, STAT, CREME96 OPEN
      DATA INUNIT/4/
      IF (NHEADER.LE.O) RETURN
      CALL CHECK_CREME96_VERSION(INFILE, IVER)
       OPEN (UNIT=INUNIT, FILE='USER:'//INFILE,
С
            STATUS='OLD', READONLY, SHARED)
С
      stat = creme96_open(infile,'user',inunit,'old')
      IF (IVER.LT.102) THEN
      DO J=1, NHEADER
         READ(INUNIT, 999) ILINE
         WRITE (OUTUNIT, 999) ILINE
 999
         FORMAT (A80)
      ENDDO
      ELSEIF (IVER.GE.102) THEN
      READ(INUNIT, 999) ILINE
      DO J=1, NHEADER
         READ(INUNIT, 999) ILINE
         WRITE (OUTUNIT, 999) ILINE
      ENDDO
      ENDIF
      CLOSE (INUNIT)
      RETURN
      END
```

```
VERSION NUMBER, PROGRAM CODE,
                                   DOSE PER SECOND,
         &
                                   ACCUMULATED DOSE)
   C
   C
          AJT 12/1/97:
   C
          Inputs: L = number of bins in integral LET spectrum
   С
                   LETMIN, LETMAX = lower, upper boundaries of LET (in MeV-cm2/g)
   С
                   LETFLUX = integral particle flux (m2-s-sr)**-1
   С
                   MODEL TYPE = I*4 label of CREME96 environment type.
          Outputs: VERSION NUMBER: CREME96 Version Number
   C
   C
                   PROGRAM CODE:
                                    DOSE program code
   C
                   DOSE PER SECOND: Dose rate (rads/second)
   C
                   ACCUMULATED DOSE: krad/sec or krad, depending upon model type
          IMPLICIT NONE
          INTEGER*4 L, MAXSPEC
          REAL*4 LETMIN, LETMAX, LETFLUX, LETVAL
          DIMENSION LETFLUX(1)
          PARAMETER (MAXSPEC=5000)
          DIMENSION LETVAL (MAXSPEC)
          INTEGER*4 VERSION NUMBER, PROGRAM CODE, I, MODEL TYPE
          REAL*4 DOSE PER SECOND, ACCUMULATED DOSE
          REAL*4 DRAD, STP, DELTA, FOURPI
   C
         WRITE (6,9998)
    9998 FORMAT(1x,' DOSE DRIVER calculation started.',
                     Please stand by.')
         DOSE PER SECOND=0.0
         ACCUMULATED DOSE=0.0
200
          CALL GET_CREME96_VERSION (VERSION NUMBER)
         PROGRAM CODE=11
         First evaluate corresponding LET Values
         IF (L.GT.MAXSPEC) THEN
              WRITE(6,995) L, MAXSPEC
    995
              FORMAT('@ 11001 ABNORMAL TERMINATION: ',
                   /,1x,' ERROR in CREME96 DOSE: ',
                   /,1x,' TOO MANY BINS: ',18,' > ',18,' max.',
        &
                   /,1x,' STOP.')
              STOP
         ENDIF
         LETVAL (1) = LETMIN
         LETVAL (L) = LETMAX
         DO 100 I=2,L-1
            LETVAL(I) = LETMIN* (LETMAX/LETMIN) ** (FLOAT(I-1)/FLOAT(L-1))
    100 CONTINUE
         Now do dose calculation
         DRAD=0.0
         DO 200 I=2,L
            STP=SQRT(LETVAL(I)*LETVAL(I-1))
           DELTA=STP*(LETFLUX(I-1)-LETFLUX(I))
           DRAD=DRAD+DELTA
    200 CONTINUE
```

SUBROUTINE CREME96 DOSE (L, LETMIN, LETMAX, LETFLUX, MODEL TYPE,

```
C
         Convert units:
         (/m2-s-sr)*(MeV-cm2/g) to rad/sec
   C
         1 \text{ rad} = 6.24E7 \text{ MeV/g}
         FOURPI=16.0*ATAN(1.0)
         DOSE_PER_SECOND=DRAD*FOURPI/6.24E11
   C
         Now calculate ACCUMULATED DOSE:
         For solar-quiet models, convert to rad/year
         IF (MODEL TYPE.EQ.0) THEN
   C
             Calculate annual dose rate (krad/yr):
           ACCUMULATED_DOSE=DOSE_PER_SECOND*31557.6
         ELSEIF (MODEL TYPE.EQ.2) THEN
             Worst-week accumulated dose (krad; in 180 hours):
   C
             ACCUMULATED DOSE=DOSE PER SECOND*648.
         ELSEIF (MODEL TYPE.EQ.1) THEN
   C
             Worst-day accumulated dose (krad; in 18 hours):
             ACCUMULATED DOSE=DOSE_PER_SECOND*64.8
         ELSEIF (MODEL TYPE.EQ.3) THEN
   C
             Peak Solar Dose rate (krad/sec):
             ACCUMULATED DOSE=DOSE PER_SECOND/1000.
         ENDIF
         WRITE(6,9999)
    9999 FORMAT(1x,' DOSE DRIVER calculation completed. Thank you.')
         IF (MODEL TYPE.EQ.0) WRITE (6,9000) DOSE PER SECOND, ACCUMULATED DOSE
ij
IF (MODEL TYPE.EQ.1) WRITE (6,9001) DOSE PER SECOND, ACCUMULATED DOSE
         IF (MODEL TYPE.EQ.2) WRITE (6,9002) DOSE PER SECOND, ACCUMULATED DOSE
IF (MODEL_TYPE.EQ.3) WRITE (6,9003) DOSE_PER SECOND, ACCUMULATED DOSE
    9000 FORMAT(' Average Dose = ',1PE13.6,' rad/sec = ',1PE13.6,
        % ' krad/year')
9001 FORMAT(' Worst-day average dose rate = ',1PE13.6,' rad/sec',
Ħ
              /,' Event-Accumulated Dose = ',1PE13.6,
į.
               ' krad in 18.0 hours.')
    9002 FORMAT(' Worst-week average dose rate = ',1PE13.6,' rad/sec',
              /,' Event-Accumulated Dose = ',1PE13.6,
' krad in 180.0 hours.')
    9003 FORMAT(' Peak SEP dose rate = ',1PE13.6,' rad/sec = ',
                  1PE13.6,' krad/sec')
         RETURN
```

END

```
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```

```
SUBROUTINE CREME96_FLUX(IZLO, IZHI, ELOWER, EUPPER, YEAR, IMODE, ITRANS,
                                GTRANSFILE, TRAPDFILE,
                                VERSION NUMBER, PROGRAM CODE,
                                M, E, FLX)
C
C
      Routine for generating the CREME96 particle fluxes.
C
      Modified 7/29/96 to output version number & program code
C
      IMPLICIT NONE
      INTEGER*4 MARR, NELM
      PARAMETER (MARR=5000, NELM=92)
      REAL*4 E,FLX
      DIMENSION E (MARR), FLX (NELM, MARR)
      REAL*4 ENERGY, DE, GET CREME96 FLUX
      INTEGER*4 IZLO, IZHI, IMODE, ITRANS, J, K, M
      INTEGER*4 VERSION NUMBER, PROGRAM CODE
      REAL*4 ELOWER, EUPPER, YEAR
      CHARACTER*80 GTRANSFILE, TRAPDFILE
      WRITE(6,9998)
 9998 FORMAT(1x,' FLUX_DRIVER calculation started. Please stand by.')
      CALL GET CREME96 VERSION (VERSION NUMBER)
      PROGRAM_CODE=3
C
      Compute energies on logaritmically-spaced grid
      DE = (EUPPER/ELOWER) ** (1./(M-1.))
      E(1)=ELOWER
      DO J=2, M-1
       E(J) = E(J-1) *DE
      END DO
      E(M)=EUPPER
      IF (ITRANS.GT.0) CALL LOAD GTF(GTRANSFILE)
      IF (ITRANS.EQ.2) CALL LOAD TRAPPED PROTONS (TRAPDFILE)
      IF (ITRANS.GT.0 .and. IMODE.GE.1) CALL LOAD SEP QSTATES
С
      Compute fluxes for each element
      DO J=IZLO, IZHI
        DO K=1, M
          ENERGY=E(K)
          FLX (J, K) = GET CREME96 FLUX (J, ENERGY, YEAR, IMODE, ITRANS)
        END DO
      END DO
      WRITE (6,9999)
 9999 FORMAT(1x,' FLUX_DRIVER calculation completed. Thank you.',
     & /,lx,' All fluxes are in units of particles/m2-s-sr-MeV/nuc.',
            ' vs. energy in MeV/nuc.',
     & /,1x,' Recommended next step:',
            ' TRANS (RUN CREME96:TRANSPORT_DRIVER)')
      RETURN
      END
```

```
Logical function creme96_inquire(filename,path)
                        The non-fully specified name of the target file.
        FILENAME:
C
С
                        Contains the VMS logical or DOS environment variable
        PATH:
C
                        pointing to file location
С
C
        Calling example:
С
C
                STAT = creme96 inquire('input.dat','creme96')
C
С
        A return value of .TRUE. indicates that the target file was
C
        found in the specified directory. .FALSE. otherwise.
С
        IMPLICIT NONE
                        file,creme96_full_filename
        character*80
        character*(*)
                        filename
        character*(*)
                        path
                        exist
        logical
        file = creme96_full_filename(filename,path)
        write(*,*)'In Inquire... fullname: ',file
        inquire(file=file,exist=exist)
        write(*,*)'Exist: ',exist
        creme96_inquire = exist
        return
        end
```

```
SUBROUTINE CREME96_LETSPEC(LETMINMG, LETMAXMG, TARGET,
                                     ELOWER, EUPPER, M, IZLO, IZUP,
        &
                                     INPUT FLUX,
        &
                                     VERSION NUMBER, PROGRAM CODE, IDIFSPEC,
        &
                                     LETMIN, LETMAX, L, SPECT, DIFSPEC)
   C
         IMPLICIT NONE
         REAL*4 LETMINMG, LETMAXMG, ELOWER, EUPPER
         REAL*4 LETMIN, LETMAX
         INTEGER*4 M, IZLO, IZUP, L
         CHARACTER*12 TARGET
         INTEGER*4 MARR, NELM, LARR
         PARAMETER (MARR=5000, NELM=92, LARR=1002)
         REAL*4 INPUT_FLUX (NELM, MARR), SPECT (LARR), DIFSPEC (LARR)
         INTEGER*4 VERSION_NUMBER, PROGRAM_CODE, IDIFSPEC
   C
         WRITE (6,9998)
    9998 FORMAT(1x,' LETSPEC DRIVER calculation started.',
                   ' Please stand by.')
         CALL GET_CREME96_VERSION(VERSION_NUMBER)
         PROGRAM_CODE=5
   C
   С
         Prepare for ULET/LETSPEC calculation.
   C
         Change units of LET range from /mg to /g
         LETMIN=1000.0*LETMINMG
         LETMAX=1000.0*LETMAXMG
  C
        Specifiy number of points in integral LET spectrum
ğasış.
E
   С
         Now calculate integral LET spectrum:
         CALL ULET96 (LETMIN, LETMAX, TARGET,
                       ELOWER, EUPPER, M, IZLO, IZUP,
                       INPUT FLUX, L, SPECT)
        &
   С
   С
   C
         Now calculate differential LET spectrum
   C
         IF (IDIFSPEC.EQ.1)
         &CALL MAKE_DIFLET_SPECTRUM(LETMIN, LETMAX, L, SPECT, DIFSPEC)
         WRITE(6,9999)
     9999 FORMAT(1x,' LETSPEC_DRIVER calculation completed. Thank you.',
         & /,1x,' Integral flux is in units of particles/m2-s-sr vs. LET',
               ' in MeV-cm2/gram.',
         & /,1x,' Recommended next step: HUP',
                ' (RUN CREME96:HI UPSET_DRIVER)')
         IF (IDIFSPEC.EQ.1) WRITE(6,9997)
     9997 FORMAT(1x,' Differential LET spectrum of flux '
              ' (in particles/m2-s-sr-(MeV-cm2/gram))',
              /,1x,' vs. LET (in MeV-cm2/gram) also calculated.')
          RETURN
          END
```

```
SUBROUTINE CREME96 TRANSPORT (INPUT FLUX,
     &
                                   ELOWER, EUPPER, M, IZLO, IZUP,
                                   IPATH, UPATHO, TARGET, SHIELDFILE,
     æ
                                   VERSION NUMBER, PROGRAM CODE,
     &
                                   OUTPUT FLUX)
C
This subroutine transports an input particle environment through a
C
    specified thickness and type of shielding. It takes account both
C
    ionization energy loss (dE/dx) as well as energy-dependent nuclear
C
    fragmentation. The output is the particle environment (differential
C
    fluxes vs. energy) inside the spacecraft, that is, 'behind' the specified
C
    shielding. This routine includes many refinements over the old CREME
C
    transport routine ("INSIDE"). Specifically:
С
C
    CREME96 TRANSPORT keeps track of projectile fragments; the old CREME
С
С
    code ignored them. This routine also uses improved Silberberg, Tsao,
    and Barghouty energy-dependent fragmentation cross-sections. Both of
С
С
    these improvements can be important for thick shielding.
С
C
    At present CREME96 TRANSPORT only does aluminum shielding; future
    versions will also offer transport through other shielding materials.
C
С
    CREME96 TRANSPORT is based on the "UPROP" code, as originally developed
C
    by John R. Letaw of Severn Communication Corp. under contract to
C
    the Gamma & Cosmic Ray Astrophysics Branch of Naval Research Laboratory
С
    in 1989. Significant improvements and "bug-extermination" have been
C
C
    provided by A.F. Barghouty of Roanoke College.
C
      IMPLICIT NONE
      CHARACTER*12 TARGET
      CHARACTER*80 SHIELDFILE
      INTEGER*4 MARR, NELM
      PARAMETER (MARR=5000, NELM=92)
      REAL*4 INPUT FLUX (NELM, MARR), OUTPUT FLUX (NELM, MARR)
      REAL*4 TEMP INPUT (NELM, MARR), TEMP FLUX (NELM, MARR)
      REAL*4 ELOWER, EUPPER, UPATHO, PATH, PSTEP, PSTEPMIN, PSTEPMAX
      REAL*4 PATHOLD, DELTA PATH, TEMP PATH
      INTEGER*4 M, N, NSP, IZLO, IZUP, IPATH, IULABEL
      REAL*4 UPATH, UUPATH, FRACSHLD
      INTEGER*4 VERSION NUMBER, PROGRAM CODE
      INTEGER*4 MAXSHIELD, NSHIELD, K, IELM, IARR
      PARAMETER (MAXSHIELD=500)
      DIMENSION UPATH (MAXSHIELD), FRACSHLD (MAXSHIELD)
      CHARACTER*5 UNITS LABEL
      DIMENSION UNITS LABEL (4)
      DATA UNITS LABEL/'g/cm2', 'mils ', 'cm ', '!!!!!'/
С
C
      WRITE(6,9998)
 9998 FORMAT(1x,' TRANSPORT_DRIVER calculation started.',
                ' Please stand by.')
      CALL GET CREME96 VERSION (VERSION NUMBER)
      PROGRAM CODE=4
C
С
      Now set parametes for transport calculation.
```

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```

```
Use recommended default:
С
      Use energy-dependent fragmentation cross-sections
С
      Use straight-ahead approximation; ignore energy spread of target fragments
C
      (This takes a lot of time and generally has only very small effect.)
C
      NSP=0
C
      Set maximum & minimum PSTEP sizes allowed in transport
C
      PSTEPMIN=0.20
      PSTEPMAX=0.20
      IF (UPATHO.GT.O.O) THEN
          NSHIELD=1
          UPATH(1)=UPATH0
          FRACSHLD(1) = 1.00
      ELSE
          CALL UNLOAD SHIELDFILE (SHIELDFILE,
                                   IPATH, NSHIELD, UPATH, FRACSHLD)
      ENDIF
      IULABEL=IPATH+1
      IF (IULABEL.GT.4) IULABEL=4
      PATHOLD=0.0
C
      DO 1000 K=1, NSHIELD
      WRITE(6,999) K, UPATH(K), UNITS LABEL(IULABEL), FRACSHLD(K)
 999 FORMAT(1x,' SHIELDING BIN ', I4,' THICKNESS = ',F10.4,1x,A5,
                   FRACTION = ', F8.4)
      UUPATH=UPATH(K)
      Get shielding thickness (PATH) in g/cm2 and transport step size:
C
      CALL UNLOAD PATH (IPATH, UUPATH, TARGET, PATH, PSTEPMIN, PSTEPMAX, PSTEP)
C
      Now perform transport:
      IF (NSHIELD.EQ.1) THEN
      CALL UPROP96 (INPUT FLUX,
                    ELOWER, EUPPER, M, IZLO, IZUP,
     æ
                    N, NSP, PATH, PSTEP, TARGET,
                    OUTPUT FLUX)
     æ
C
C
      ELSE
C
C
      Modification 8-16-96 by AJT:
C
      To speed up calculations through thick shielding distributions,
C
      allow output of one step to be input to the next step.
      DELTA PATH=PATH-PATHOLD
      IF (DELTA PATH.LT.O.) DELTA PATH=0.0
      DO 300 IELM=1, NELM
         DO 200 IARR=1, MARR
                 IF (K.EQ.1) THEN
                 TEMP INPUT (IELM, IARR) = INPUT FLUX (IELM, IARR)
                 TEMP PATH=PATH
                 ELSEIF (K.GT.1 .and. DELTA PATH.LT.PSTEP) THEN
                 TEMP INPUT(IELM, IARR) = INPUT_FLUX(IELM, IARR)
                 TEMP PATH=PATH
```

```
ELSEIF (K.GT.1 .and. DELTA PATH.GE.PSTEP) THEN
                    TEMP_INPUT(IELM, IARR) = TEMP_FLUX(IELM, IARR)
                    TEMP_PATH=DELTA PATH
                    ENDIF
     200
            CONTINUE
     300 CONTINUE
         CALL UPROP96 (TEMP INPUT,
        &
                       ELOWER, EUPPER, M, IZLO, IZUP,
        &
                       N, NSP, TEMP PATH, PSTEP, TARGET,
                       TEMP_FLUX)
         PATHOLD=PATH
   C
         Now add to weighted sum:
         DO 500 IELM=1, NELM
            DO 400 IARR=1, MARR
               OUTPUT_FLUX(IELM, IARR) =OUTPUT_FLUX(IELM, IARR) +
        &
                                       TEMP FLUX (IELM, IARR) *FRACSHLD(K)
     400
            CONTINUE
     500 CONTINUE
         ENDIF
   1000 CONTINUE
         WRITE (6, 9999)
    9999 FORMAT(1x,' TRANSPORT_DRIVER calculation completed. Thank you.',
        & /,1x,' All fluxes are in units of particles/m2-s-sr-MeV/nuc',
              ' vs. energy in MeV/nuc.',
        & /,1x,' Recommended next step: ',
        & /,5x,' LETSPEC',
              ' (RUN CREME96:LETSPEC_DRIVER)'
Ŀi
             ' for heavy-ion induced SEUs;',
        & /,2x,' or PUP
                          (RUN CREME96: PROTON UPSET_DRIVER)',
' for proton-induced SEUs.')
        RETURN
         END
```

:::

```
REAL FUNCTION CRF96 (IZ, EN, YEAR, IMODE)
С
С
        THIS ROUTINE RETURNS THE DIFFERENTIAL FLUX IN PARTICLES/((M**2)*
C
        STER*SEC*MEV/U) AS IT IS FOUND IN THE INTERPLANETARY MEDIUM
C
       NEAR EARTH and OUTSIDE the magnetosphere
C
С
       IZ = ATOMIC NUMBER OF THE IONS.
С
       E = ENERGY (IN MEV/U).
С
       Y = THE YEAR: 1975.144=SOLAR MIN; 1980.598=SOLAR MAX.
C
       M = Particle environment model
С
            0 = non-solar particles only: GCR+ACR
C
            1 = "Worst day" Solar Energetic Particle Environment
С
            2 = "Worst week" Solar Energetic Particle Environment
C
            3 = "Peak (worst 5-minutes) Solar Energetic Particle Environment
C
C
        IMPLICIT NONE
        INTEGER*4 IZ, IQ, IMODE, IDUM
        REAL*4 EN, YEAR, GCRF, GCR_FLUX, ACRF, ACR_FLUX, SEP_FLUX
        CRF96=0.0
        IF (IMODE.LT.O. .or. IMODE.GT.3) RETURN
        IF (EN.LT.O.) RETURN
        IF (IZ.LT.1 .or. IZ.GT.92) RETURN
       IF (IMODE.EQ.0) THEN
            Get Galactic Cosmic Ray contribution
            GCRF=GCR_FLUX(IZ,EN,YEAR,IDUM)
            Get Anomalous Cosmic Ray contribution
            ACRF=0.0
            DO 100 IQ=1, IZ
            ACRF=ACRF+ACR_FLUX(IZ, IQ, EN, YEAR)
100
            CONTINUE
            CRF96=GCRF+ACRF
       ELSEIF (IMODE.NE.0) THEN
            CRF96=SEP_FLUX(IZ,EN,IMODE)
       ENDIF
       RETURN
        END
```

```
*******************
    C SUBROUTINE CTABLE in Module UPROP.FOR
    C
    C Creates the auxiliary spallation cross section data file (CTABLE.DAT) if
    C it does not already exist. It also calculates energy losses associated
    C with spallation reactions (Sept. 1993).
   C Modified 06-05-96: add NSP to arguments, to control PARTIALS
   C Modified 11-17-97: add IMPLICIT NONE and variable-type declarations.
   C Parameters
             Maximum atomic number of elements to be transported (<= 109)
   C NELM
             Maximum number of energies at which cross section data are
   C MCS
             defined
   C ELOWER Lower energy bound of input and output spectra (>= 0.1 MeV)
   C EUPPER Upper energy bound of input and output spectra (<= 100000 MeV)
            Number of logarithmically equally-spaced energy bins at which
   C
             cross sections are evaluated (ABS(N) < MCS)
   C NSP
            =1 turns on nuclear dE/dx in PARTIALS; 0 otherwise
   C IZLO Least atomic number of elements transported (>= 1)
   C IZUP Greatest atomic number of elements transported (<= 109)
   C TARGET Name of the target shielding material (<= 12 bytes)
   C
   C Important variables
   C ECS
            Energy at each cross section grid point.
   C CT
            Temporary array for holding the spallation cross sections of
            one element at one energy.
   C ELOSS Temporary array for holding the energy loss of one element
   C
           at one energy averaged over each fragment's isotopes.
   C TOTAL Abundance weighted elemental cross section.
            Cross section array for all elements at a single energy.
C ENLOSS Energy loss array for all elements at a single energy averaged
TU C
            each element's isotopes.
Li C
C Subprograms
  С
C SUBROUTINE MFP (ENERGY, K, ALL, TARGET, PATH)
      Returns the mean free path PATH in g/cm**2 at energy ENERGY for an
      element with charge K and mass ALL in target material TARGET
   C
   C SUBROUTINE PARTIALS (ENERGY, K, ALL, TARGET, CT, ANORM)
      Returns the partial spallation mean free paths CT in g/cm**2 at energy
  C
      ENERGY for an element with charge K and mass ALL in target material
  C
      TARGET.
  C
  C BLOCK DATA DO1
      Defines the atomic masses of elements in the range 1 <= Z <= 109 and
  C
  С
      places them in the array AMASS
  C
  C Data Files
  C
  C PERIODIC.DAT
      Contains a list of the isotopes of each element and their natural
  С
  С
      abundance.
  \mathbf{c}
  C CTABLE.DAT
      Contains nuclear spallation cross section data for the transport
```

SUBROUTINE CTABLE (ELOWER, EUPPER, N, NSP, IZLO, IZUP, TARGET)

```
calculation. Automatically created by this subroutine when needed.
   IMPLICIT NONE
         INTEGER NELM, MCS
         PARAMETER (NELM=92, MCS=10)
         REAL*4 CABU (NELM, 9), TOTAL (NELM)
         REAL*4 ECS (MCS), C (NELM, NELM), CT (NELM), NORM (NELM)
         REAL*4 CS (NELM, NELM), ENLOSS (NELM, NELM), ELOSS (NELM), ETOTL (NELM)
         INTEGER*4 CISO(NELM, 9), STAT, CREME96_OPEN
         CHARACTER*12 TARGET, TARGET$
         REAL*4 AMASS
         COMMON/MASS/AMASS(109)
         REAL*4 ELOWER$, EUPPER$, ELOWER, EUPPER
         INTEGER*4 N$, IZLO$, IZUP$, NSP$, N, IZLO, IZUP, NSP
         INTEGER*4 I, J, K, L
         REAL*4 ENERGY, DENERGY, ALL, ANORM, PATH, REALNORM, FPRO, FALP
         DATA ELOWER$, EUPPER$, N$, IZLO$, IZUP$, NSP$/0.,0.,0,0,0/
         DATA TARGET$/' '/
   C
         FORMAT Statements
  C Format statement modified 6-5-96 by AJT to accomodate NSP
  100
         FORMAT (1X, 2 (1PE10.4, 2X), 4 (15, 2X), A12, 2X, 1PE10.4)
   200
         FORMAT ((1X,6(1PE11.4,2X)))
   300
         FORMAT (6X, 9(1X, I3, 1X, F4.2))
C
         Otherwise perform calculation of cross sections
   С
         Read in list of elements from PERIODIC.DAT
L
差
         OPEN (UNIT=15, FILE='CREME96: PERIODIC.DAT', STATUS='OLD',
               READONLY, SHARED)
T.
         stat = creme96 open('periodic.dat','cr96tables',15,'old')
E.
         DO J=1,83
<u>Lik</u>
           READ (15,300) (CISO(J,K),CABU(J,K),K=1,9)
END DO
         CLOSE (UNIT=15)
   C
         Open output data file
          OPEN(UNIT=13, FILE='USER: CTABLE.DAT', STATUS='NEW')
          stat = creme96_open('ctable.dat','user',13,'new')
          OPEN(UNIT=17,FILE='USER:SPTABLE.DAT',STATUS='NEW')
          stat = creme96 open('sptable.dat','user',17,'new')
  C
         Write header
         ELOWER$=ELOWER
         EUPPER$=EUPPER
        N$=N
         IZLO$=IZLO
         IZUP$=IZUP
        NSP$=NSP
         TARGET$=TARGET
        WRITE (13,100) ELOWER, EUPPER, N, IZLO, IZUP, NSP, TARGET
         WRITE (13, '(A)') ''
         WRITE (17,100) ELOWER, EUPPER, N, IZLO, IZUP, NSP, TARGET
```

```
C
       Compute vector of energies
       IF (N.GE.2) THEN
         ECS(1)=ELOWER
         DENERGY= (EUPPER/ELOWER) ** (1./FLOAT(N-1))
         DO J=2,N
           ECS(J) = ECS(J-1) *DENERGY
         END DO
       ELSE
         ECS(1) = 2000.
       ENDIF
C
       Compute parameters
       DO J=1,N
         ENERGY=ECS (J)
C
       Initialize some arrays
         DO K=1, NELM
           TOTAL(K) = 0.
           NORM(K) = 0.
           DO I=1, NELM
             C(K,I)=0.
             ENLOSS(K, I) = 0.
             CS(K,I)=0.
           END DO
         END DO
        DO K=IZLO, IZUP
С
      For each incident particle at each energy compute
С
      partial cross sections, total cross section, and the
      normalization factor. Average over isotopes.
          DO L=1,9
             IF (CABU(K,L).GT.O.) THEN
             ALL=REAL(CISO(K,L))
             CALL PARTIALS (ENERGY, K, ALL, TARGET, CT, NSP, ELOSS, ANORM)
             CALL MFP (ENERGY, K, ALL, TARGET, PATH)
             DO I=2,K+1
               C(I,K) = C(I,K) + CT(I) * CABU(K,L)
               CS(I,K) = C(I,K) !Cross section w/o merging!
             END DO
             DO I=1, K+1
                           !Energy-loss averaged over isotopes of K:
               ENLOSS(I,K) = ENLOSS(I,K) + ELOSS(I) * CABU(K,L)
             END DO
             NORM(K) = NORM(K) + ANORM*CABU(K, L)
             TOTAL (K) = TOTAL (K) + PATH * CABU (K, L)
             ENDIF
          END DO
C
      Allow for further renormalization
      REALNORM=1.
      Compute cross sections for proton and alpha production
С
      The procedure used here is taken from J.R. Letaw,
```

WRITE (17, '(A)') ' '

Œ

į.

r.

į.i.

C

Phys. Rev. C28, 2178 (1983).

```
IF (K.GT.2) THEN
   C
                  FF=AMASS(K)/FLOAT(K)+0.67
   C
                  ASSUME 15% He AND 85% H in product
   C
                  FPRO=1./(1.+2a/p)
                  FPRO=0.739
   C
                  FALP=1./(2.+ p/a)
                  FALP=0.130
                  C(1,K)=FPRO*(TOTAL(K)*FLOAT(K)-NORM(K)*REALNORM)
                  C(2, K) = FALP * (TOTAL(K) * FLOAT(K) - NORM(K) * REALNORM) + C(2, K)
               ENDIF
   С
          Compute partials for alpha or proton into proton
               IF (K.EQ.1) C(1,K)=0.
               IF (K.EQ.2) C(1,K)=2.0*TOTAL(K)
   С
                CS(1, K) = C(1, K)
   С
                CS(2,K) = C(2,K)
   C
   С
          Merge total cross sections in
               C(K,K) = C(K,K) - TOTAL(K)
               CS(K,K) = CS(K,K) + TOTAL(K)
   Ċ
          Output results for current energy
□ C
            END DO
            \mathtt{WRITE}\,(13\,,200)\quad(\,(\mathtt{C}\,(\mathtt{K}\,,\mathtt{I})\,\,,\mathtt{K=IZLO}\,,\mathtt{IZUP})\,\,,\mathtt{I=IZLO}\,,\mathtt{IZUP})
Nuclear stopping power table:
                                               Sept. 1993
            DO K=IZLO, IZUP
                ETOTL (K) = 0.
i.
                DO I=IZLO, IZUP
                  ETOTL(K) = ETOTL(K) + ENLOSS(K, I) *CS(K, I)
į.
                END DO
FL
            END DO
Ŀ
            WRITE(17,200) (ETOTL(K), K=IZLO, IZUP)
          END DO
  C
  C
          Close output file and stop
          CLOSE (UNIT=13)
          CLOSE (UNIT=17)
          RETURN
          END
```

```
SUBROUTINE SPLINE ( X, Y, N, NMAX, NATURAL, YP1, YPN, Y2, U )
   C
   С
          Given arrays X and Y of length N containing a tabulated function
          Y(i) = f[X(i)] specified upon an set of montonically increasing
   C
   C
          arguments \{X(1) < X(2) < ... < X(N)\}, SPLINE produces the array Y2
          containing the second derivative of the interpolating function at
   C
          the same arguments. The boundary conditions are specified by the
   С
   С
         logical array
   С
   С
                  NATURAL(1) = .TRUE.
                                       : Y2(1) = 0.0
   С
                  NATURAL(1) = .FALSE. : Y2(1) = YP1
   C
   С
                  NATURAL(2) = .TRUE. : Y2(N) = 0.0
   C
                  NATURAL(2) = .FALSE. : Y2(N) = YPN.
   C
         Note that dummy arguments YP1 and YPN must be supplied even if
   С
   C
         not used.
   С
   C
         The algorithm uses a scratch array U, which is included in the
         argument list so that its dimensions need not be adjusted for
   C
   C
         each application. The maximum dimension for the input and
   C
         output arrays is NMAX:
   C
   С
         Adapted from "Numerical Recipes", by W.H. Press et al.
  C
           IMPLICIT REAL*8 (A-H, O-Z)
J
  cx
IMPLICIT REAL*4 (A-H, O-Z) !for use with minfun
         LOGICAL NATURAL (2)
         DIMENSION X (NMAX), Y (NMAX), Y2 (NMAX), U (NMAX)
FL.
С
          ... Check arguments and boundary conditions.
   C
IF (N .GT. NMAX) STOP ' Too many points in SPLINE.'
         IF ( NATURAL(1) ) THEN
TU
           Y2(1) = 0.0
IJ
           U(1) = 0.0
ļ.
         ELSE
Y2(1) = -0.5
           U (1) = (3.0 / (X(2) - X(1))) * ((Y(2) - Y(1)) /
                           (X(2) - X(1)) - YP1)
         END IF
   C
         IF ( NATURAL(2) ) THEN
           QN = 0.0
           UN = 0.0
           Y2(N) = 0.0
           U(N) = 0.0
         ELSE
           QN = 0.5
           UN = (3.0/(X(N) - X(N-1)))*(YPN - (Y(N) - Y(N-1))/
                     (X(N) - X(N-1))
        END IF
  C
  C
         ... Decomposition loop of tridiagonal algorithm. Y2 and U are used
  C
             for temporary storage of decomposed factors.
  C
        DO I = 2, N-1
           SIG
                 = (X(I) - X(I-1)) / (X(I+1) - X(I-1))
                 = SIG * Y2(I-1) + 2.0
```

```
Y2(I) = (SIG - 1.0) / P
         U(I) = (6.0 * ((Y(I+1) - Y(I) ) / (X(I+1) - X(I) ) -
                         (Y(I)
                                - Y(I-1)) / (X(I)
                                                    - X(I-1))) /
                         (X(I+1) - X(I-1)) - SIG*U(I-1)) / P
       END DO
       Y2(N) = (UN - QN*U(N-1)) / (QN*Y2(N-1) + 1.0)
 С
 C
       ... Backsubstitution loop.
 C
       DO K = N-1, 1, -1
        Y2(K) = Y2(K)*Y2(K+1) + U(K)
       END DO
 C
       RETURN
       END
       SUBROUTINE SPLINT ( XA, YA, Y2A, N, X, Y )
 С
      Given arrays XA and YA of length N, which tabulate a function
С
       (with the XA(i)'s in order), and given the array Y2A, which is
С
      output from SPLINE above, and given a value of X, this routine
С
       returns a cubic-spline interpolated value Y.
С
С
С
      Adapted from "Numerical Recipes", by W.H. Press et al.
C
CX
        IMPLICIT REAL*8 (A-H, O-Z)
      IMPLICIT REAL*4 (A-H, O-Z) !for use with minfun
      DIMENSION XA(N), YA(N), Y2A(N)
C
C
       ... Locate nearest base points by bisection.
      KLO = 1
      KHI = N
    1 IF ( (KHI - KLO) .GT. 1) THEN
        K = (KHI + KLO) / 2
        IF (XA(K) .GT. X) THEN
          KHI = K
        ELSE
          KLO = K
        END IF
        GO TO 1
      END IF
      H = XA(KHI) - XA(KLO)
      IF (H .EQ. 0.0) STOP ' Arguments for SPLINE must be unique.'
C
      ... Evaluate cubic spline polynomial.
С
C
      A = (XA(KHI) - X
      B = (X
                   - XA(KLO)) / H
      Y = A*YA(KLO) + B*YA(KHI)
     . + ( A*(A*A - 1)*Y2A(KLO) + B*(B*B - 1)*Y2A(KHI) )*H*H / 6.0
C
     RETURN
     END
```

```
REAL FUNCTION CUT96 (IZ, EN, YEAR, IMODE)
C
C
        THIS ROUTINE OBTAINS DIFFERENTIAL PARTICLE FLUXES AND
С
        APPLIES THE GEOMAGNETIC CUTOFF TRANSMISSION FUNCTION
C
        AND RETURNS THE RESULTING FLUX, MODULATED TO THE
C
        ORBIT-AVERAGE CUTOFF.
С
C
        IZ = ION ATOMIC NUMBER.
C
        EN = ION ENERGY IN MEV/AMU.
        YEAR = YEAR (1975.144 = SOLAR MIN.; 1980.598 = SOLAR MAX.).
C
C
        IMPLICIT NONE
        INTEGER*4 IZ, IQ, IMODE, KZ, IDUM, J
        REAL*4 EN, YEAR, A, SEP QSTATES, AN, Q, P, TRF
        REAL*4 MAGNETIC_RIGIDITY, GET_GTF, GCR_FLUX, ACR_FLUX, SEP_FLUX
        REAL*4 GCRF, ACRF, ACRFQ, SF
        COMMON/MASS/A(109)
        COMMON/SEP_QSTATES/SEP_QSTATES(30,30)
        CUT96=0.0
        IF (IMODE.LT.O .or. IMODE.GT.3) RETURN
        IF (EN.LT.O.) RETURN
        IF (IZ.LT.1 .or. IZ.GT.92) RETURN
        AN=A(IZ)
        IF (IMODE.EQ.0) THEN
            Galactic-Cosmic Ray Component
            Q=IZ*1.0
            P=MAGNETIC RIGIDITY (EN, Q, AN)
            TRF=GET_GTF(P)
            GCRF=GCR_FLUX(IZ, EN, YEAR, IDUM)
            CUT96=CUT96+GCRF*TRF
            Anomalous Component
            ACRF=0.0
            DO 100 IQ=1,IZ
            ACRFQ=ACR_FLUX(IZ, IQ, EN, YEAR)
            IF (ACRFQ.GT.0) THEN
               Q=IQ*1.0
               P=MAGNETIC RIGIDITY (EN, Q, AN)
               TRF=GET GTF(P)
               ACRF=ACRF+ACRFQ*TRF
            ENDIF
  100
            CONTINUE
            CUT96=CUT96+ACRF
        ELSEIF (IMODE.NE.0) THEN
            Solar Energetic Particle Contribution
C
            SF=SEP FLUX(IZ, EN, IMODE)
            IF (SF.EQ.O.) RETURN
            KZ=IZ
C
            For elements heavier than Zn, use Zn charge states
            IF (KZ.GT.30) KZ=30
```

```
DO 3200 J=1, KZ

TRF=0.0

IF (SEP_QSTATES(KZ,J).GT. 1.0E-8) THEN

Q=J*1.0

P=MAGNETIC_RIGIDITY(EN,Q,AN)

TRF=GET_GTF(P)

ENDIF

CUT96=CUT96+SF*SEP_QSTATES(KZ,J)*TRF

CONTINUE

ENDIF
```

RETURN END END

```
BLOCK DATA DO1
     *******************
     Atomic Mass Tabulation from Review of Particle Properties
C
С
                               Physics Letters B204 (April, 1988)
COMMON/MASS/AMASS(109)
     DATA AMASS/
    & 1.00794,4.002602,6.941,9.012182,10.811,12.011,14.00674,15.9994,
    & 18.9984032,20.1797,22.989768,24.305,26.981539,28.0855,30.973762,
    & 32.066,35.4527,39.948,39.0983,40.078,44.95591,47.88,50.9415,
    & 51.9961,54.93805,55.847,58.9332,58.69,63.546,65.39,69.723,
    & 72.61,74.92159,78.96,79.904,83.80,85.4678,87.62,88.90585,91.224,
    & 92.90638,95.94,98.,101.07,102.9055,106.42,107.8682,112.411,
    & 114.82,118.71,121.75,127.6,126.90447,131.29,132.90543,137.327,
    & 138.9055,140.115,140.90765,144.24,145,150.36,151.965,157.25,
    & 158.92534,162.5,164.93032,167.26,168.93421,173.04,174.967,
    & 178.49,180.9479,183.85,186.207,190.2,192.22,195.08,196.96654,
    & 200.59,204.3833,207.2,208.98037,209.,210.,222.,223.,226.0254,
    & 227.0278,232.0381,231.03588,238.0289,237.0482,244.,243.,247.,
    & 247.,251.,252.,257.,258.,259.,260.,261.,262.,263.,262.,265,266/
```

```
REAL*4 FUNCTION DIFPLD(S,L,W,H)
```

```
C
   С
                 THIS FUNCTION RETURNS THE PROBABILITY DENSITY FOR PATHS
   С
           OF LENGTH S THROUGH A PARALLELEPIPED OF DIMENSIONS
   C
           L, W, AND H. S, L, W, AND H MUST BE IN THE SAME UNITS.
   C
                 THIS IS AN EXACT SOLUTION, DUE TO M. D. PETROFF OF
   C
           ROCKWELL INTERNATIONAL (SEE J. C. PICKEL AND J. T. BLANDFORD,
   C
           IEEE TRANS. ON NUCL. SCI. NS-27, 1006(1980)) WITH
   С
           SIMPLIFICATIONS DUE TO WARREN BENDEL OF NRL (PRIVATE
   С
           COMMUNICATION). THE EQUATION NUMBERS REFER TO THE APPENDIX
   C
           OF PICKEL AND BLANDFORD'S PAPER.
   C
   C
           Modified by AJT 4-2-96: IMPLICIT NONE and variable-type declarations
   C
           added
   C
           IMPLICIT NONE
           REAL*4 S, L, H, W, AP, G
   C
   C
           EQUATION (A-7)
   C
           AP=3.*(H*W+H*L+L*W)
  C
   C
           EQUATION (A-8)
  C
           DIFPLD = (G(S, L, W, H) + G(S, W, L, H) + G(S, L, H, W) + G(S, W, H, L) +
1 G(S,H,W,L)+G(S,H,L,W))/(3.1416*AP)
           RETURN
           END
REAL*4 FUNCTION G(S,X,Y,Z)
IMPLICIT NONE
REAL*4 S,X,Y,Z,KSQ,T,RSQ,R,V,PSQ,QSQ,TSQ
<sup>®</sup> C
E C
           PRELIMINARY DEFINITIONS
TU c
IJ
           KSQ=X*X+Y*Y
Ŀ
           TSQ=X*X+Z*Z
ų)
           T=SQRT (TSQ)
           RSQ=KSQ+Z*Z
           R=SQRT (RSQ)
           V=12.*X*Y*Z*Z
           PSQ=S*S-Z*Z
           QSQ=S*S-X*X-Z*Z
           IF((S.GE.0.0).AND.(S.LT.Z)) GO TO 10
           IF((S.GE.Z) .AND.(S.LT.T))GO TO 20
           IF((S.GE.T) .AND.(S.LE.R))GO TO 30
          G=0.0
          RETURN
  C
  C
          EQUATION (A-9)
  C
   10
          G=8.*Y*Y*Z/KSQ-S*(3.*X*Y/(R*T))**2
          RETURN
  C
  C
          EQUATION (A-10)
  C
   20
          G=S*(3.*Y/SQRT(KSQ))**2-S*(3.*X*Y/(T*R))**2
       1 -X*(SQRT(PSQ)/S)*(8.+4.*Z*Z/(S*S))
       2 + (V*ATAN(Y/X) - (Y*Z*Z/SQRT(KSQ))**2)/(S*S*S)
```

```
PROGRAM DOSE DRIVER
         IMPLICIT NONE
         CHARACTER*80 INFILE, OUTFILE
         REAL*4 EMINCUT, EMAXCUT, ELOWER, EUPPER
         INTEGER*4 IZMIN, IZMAX, IZLO, IZUP, M, L
         CHARACTER*12 TARGET
         INTEGER*4 MARR, NELM, LARR
         PARAMETER (MARR=5000, NELM=92, LARR=1002)
         REAL*4 INPUT FLUX (NELM, MARR), LETFLUX (LARR)
         INTEGER*4 VERSION NUMBER, PROGRAM CODE
         REAL*4 LETMINMG, LETMAXMG, LETMIN, LETMAX
         REAL*4 DOSE PER_SECOND, ACCUMULATED DOSE
         INTEGER*4 MODEL TYPE
  C
  С
         Get parameters of dose calculation:
  C
        CALL INIDOSE (INFILE, LETMINMG, LETMAXMG,
                       IZMIN, IZMAX, EMINCUT, EMAXCUT,
                       TARGET, OUTFILE)
  C
        Unload input particle flux file into array:
  C
        CALL UNLOAD_PARTIAL_FLUX(INFILE, IZMIN, IZMAX, EMINCUT, EMAXCUT,
                                   ELOWER, EUPPER, M, IZLO, IZUP,
INPUT FLUX)
        Check model-type
        CALL GET_CREME96 FLUX MODEL (INFILE, MODEL TYPE)
C
        Now do integral LET spectrum calculation:
  C
        LETMIN=LETMINMG*1000.0
LETMAX=LETMAXMG*1000.0
L=LARR
incia.
I.
        CALL ULET96 (LETMIN, LETMAX, TARGET,
                     ELOWER, EUPPER, M, IZLO, IZUP,
       &
                     INPUT FLUX, L, LETFLUX)
  C
  C
        Now do numerical integration to get dose value:
  С
  С
            DOSE_PER_SECOND = average dose rate (rads/second)
  C
            ACCUMULATED_DOSE = krad or krad/sec, depending on MODEL TYPE
  С
        CALL CREME96_DOSE(L, LETMIN, LETMAX, LETFLUX, MODEL_TYPE,
       &
                           VERSION NUMBER, PROGRAM CODE,
                           DOSE PER_SECOND, ACCUMULATED_DOSE)
  C
  С
        Now write dose results to output file:
  C
```

CALL OUTPUT_CREME96_DOSE(INFILE,IZLO,IZUP,LETMIN,LETMAX,

OUTFILE)

C C EMINCUT, EMAXCUT, TARGET, MODEL_TYPE,

VERSION_NUMBER, PROGRAM_CODE,
DOSE_PER_SECOND, ACCUMULATED_DOSE,

```
&
                       (EN, NPTS, IPARAM, PARAMS, XSECT FILE, XSECT)
   C
           Subroutine to evaluate SEU cross-section for an array
   С
   C
           of abscissa values:
   C
   C
           This same routine is used for both proton-induced and heavy-ion
   C
           induced cross-sections; but the dimensions of the inputs and
   С
           output are different in the two cases.
   C
   C
   С
           INPUTS: EN:
                           array of proton energies (in MeV) for proton SEUS
   C
                        OR array of LET values (in MeV-cm2/mg) for heavy-ion SEUs
   С
                   NPTS: number of points in the array
   C
                   IPARAM: specifies cross-section model or format:
   C
                           IPARAM=0: table of values
   C
                           IPARAM=1: Bendel 1-parameter fit
   C
                           IPARAM=2: Bendel 2-parameter fit
   C
                           IPARAM=4: Weibull fit
   С
                   PARAMS: array of at least dimension IPARAM, containing
   C
                           the fit parameters for IPARAM=1,2, or 4.
   C
   C
                   XSECT FILE: name of file containing cross-section table
   C
                               (for IPARAM=0 option). Cross-section values
   C
                               will be linearly interpolated in this table,
   С
                               with zero below the first entry's abscissae
   C
                               and a plateau value at the the last entry's
   C
                               ordinate
   C
           OUTPUT: XSECT:
                             array containing the cross-section values
   C
                             corresponding to values in EN array
   C
                             in 1.0E-12 cm2/bit (for proton cross-sections)
                        OR
                             in 1.0E-8 cm2/bit (for heavy ion cross-sections)
į.
   С
   С
           Written by:
                          Allan J. Tylka
   С
                          Code 7654
                          Naval Research Laboratory
                          Washington, DC 20375-5352
Ŀ
  C
                          tylka@crs2.nrl.navy.mil
  C
  C
          Last update: 29 March 1996
   C-----
          IMPLICIT NONE
          INTEGER*4 IPARAM, K, NPTS, NSV, NSVMAX
          REAL*4 EN, XSECT, PARAMS, A, B, O, W, P, BENDEL1, BENDEL2, WEIBULL
          REAL*4 XV, YV
          REAL*4 INTERPOLATE XSECT TABLE
          CHARACTER*80 XSECT FILE
          PARAMETER (NSVMAX=5000)
          DIMENSION XV(NSVMAX), YV(NSVMAX)
          DIMENSION EN(1), XSECT(1), PARAMS(4)
          IF (IPARAM.EQ.0) THEN
              CALL UNLOAD_XSECT_FILE(XSECT_FILE, NSV, XV, YV)
          ELSEIF (IPARAM.EQ.1) THEN
              A=PARAMS(1)
          ELSEIF (IPARAM.EQ.2) THEN
```

A=PARAMS(1) B=PARAMS(2)

SUBROUTINE EVALUATE SEU CROSS SECTION

```
ELSEIF (IPARAM.EQ.4) THEN
           O=PARAMS(1)
           W=PARAMS(2)
           P=PARAMS(3)
           A=PARAMS (4)
       ELSE
           WRITE(6,9999) IPARAM
9999
           FORMAT ('@ 10001 ABNORMAL TERMINATION: ',
           /,1x,' ERROR in EVALUATE_SEU_CROSS_SECTION: ',
    &
           /,1x,' CROSS-SECTION STEERING CODE UNKNOWN: ',15,
           /,1x,' STOP.')
           STOP
       ENDIF
       IF (NPTS.LE.0) RETURN
       DO 1000 K=1, NPTS
          XSECT(K) = 0.0
          IF (IPARAM.EQ.0) THEN
              XSECT(K) = INTERPOLATE_XSECT_TABLE(NSV, XV, YV, EN(K))
          ELSEIF (IPARAM.EQ.1) THEN
              XSECT(K) = BENDEL1(A, EN(K))
          ELSEIF (IPARAM.EQ.2) THEN
              XSECT(K) = BENDEL2(A, B, EN(K))
          ELSEIF (IPARAM.EQ.4) THEN
              XSECT(K) = WEIBULL(O, W, P, A, EN(K))
          ENDIF
1000
       CONTINUE
       RETURN
       END
```

```
SUBROUTINE E_LOSS(IZ, IA, JZ, JA, KZ, KA, ELAB, dKE, SigmaKE)
           C
          С
                            Computes the average energy loss dKE and variance SigmaKE when
          C
                            nuclide (IZ, IA) impinges on medium (JZ, JA)
          C
                            producing fragment (KZ,KA).
          C
                            Fragment is no longer at energy ENERGY, i.e. straight-
          C
                            ahead approximation is relaxed. Medium can be Hydrogen,
                            Helium, or any other nuclide.
          С
                            Based on the paper by Barghouty, Tsao, and Silberberg, 23rd ICRC,
          C
          C
                            Calgary, Canada, 1993.
          C
          C
                            January 1994
          C
                            INTEGER AP, ZP, AT, ZT, AA, AAZ, BB, BBZ, CC, CCZ
                            INTEGER AF, ZF, AFF, ZFF, DELTA A, DELTA Z, AC, ZC, ATO, ZTO
         C
                           DATA WN, EFERMI / 931.504,38./
                           DATA CONST, RO, PI/1.44, 1.2, 3.14159/
                           DATA Ebin/20./
                           DATA IENT/0/
         C
        C
The state of the s
                           IF (IENT.EQ.0) THEN
                                      IENT=1
                                      WRITE (6,9999)
                                      FORMAT(1x,' In nuclear transport: Subroutine E_LOSS active.')
            9999
                           ENDIF
                           dKE=0.
                           SigmaKE=0.
Č C
=
                          IF(ELAB.LT.50.) RETURN
i c
ZP=IZ
14
                         AP=IA
i,
                          ZT=JZ
AT=JA
                          ZF=KZ
                          AF=KA
        С
                         ZFF=KZ
                         AFF=KA
        С
                         IF(AF.GE.AP) RETURN
        C
                         PLAB=SQRT((ELAB+WN)**2-WN**2)
                         Pbin=(Ebin/ELAB)*Plab
        C
       C....Energy loss calculation for Z(TARGET) < 6:</pre>
                         SCALE=1.
                         IF(ZT.LT.6) THEN
                                  SCALE = (6.+ZT)/12.
                                 AT=12
                                  ZT=6
                         END IF
       C
       C....Energy loss calculation for Z(PROJECTILE) <6:</pre>
                         SCALE=SCALE*1.
```

```
IF(ZP.LT.6) THEN
            IF (ZP.EQ.1) THEN
              dKE=0.
              DP=0.
              RETURN
            ELSE
              SCALE = SCALE * (6.+ZP)/12.
              ZP=6
            END IF
        END IF
  С
  C.... Energy loss calculation for all other nuclides:
        CALL GLBR (AP, ZP, AT, ZT, AA, AAZ, BB, BBZ, CC, CCZ)
  C
        Projectile and target A and Z numbers:
        A0=AP
        AZ0=ZP
        B0=AT
        BZ0=ZT
        A=AA
        AZ = AAZ
        B=BB
        BZ=BBZ
        C=CC
        CZ = CCZ
C C C 8
        Q=HEAT(A0,AZ0,B0,BZ0,ELAB,A,AZ,B,BZ,C,CZ,TA,TB,TC,Temp)
        Relative size of fragment to source "A":
        DELTA A=A-AF
        DELTA Z=AZ-ZF
≅ C
        Coulomb Barrier:
1
        Ec = (CONST*(AZ-1.))/(R0*(SQRT(A-1.)+1.))
filc
IF (DELTA A.GE.1) THEN
L-C
        CHECK IF FRAGMENT IS TOO SMALL TO BE A SPALLATION PRODUCT.
道C
        Here we make the assumption that if the fragment is too small,
÷.iC
        i.e., fragment size < AP/2, it is accompanied by a heavy partner.
 C
        We proceed to calculate the loss of that heavy partner assuming
 C
        further that both partners suffer the same energy loss per nucleon.
 C
 C
          IF (AF.LT.DELTA A) THEN
            AF=DELTA_A
            ZF=DELTA Z
            GO TO 8
          END IF
 C
          AFMASS=AF*WN+DROP(AF, ZF)
          AMASS=A*WN+DROP(A,AZ)
 C
          CONSERVE MASS:
          DMASS=AMASS-AFMASS
         TD = DELTA_A*(TA+ (3./5.)*EFERMI) + DELTA_Z*Ec
         CONSERVE TOTAL ENERGY:
         TF = (AMASS+TA*A) - (DMASS+TD) - AFMASS
         PF = SQRT((AFMASS+TF)**2-AFMASS**2)
         dKE=ELAB-TF/AF
         dKE=SCALE*dKE
```

```
IF(dKE.LT.0.) dKE=0.
            PERC=(dKE/ELAB) *100.
            DP=PLAB-PF/AF
         ELSE
           AC=AP+(AT-B)
           ZC=ZP+(ZT-BZ)
          DELTA A=AC-AF
          DELTA Z=ZC-ZF
         Coulomb Barrier:
   C
         Ec = (CONST*(ZC-1.))/(R0*(SQRT(AC-1.)+1.))
   C
           AFMASS=AF*WN+DROP(AF, ZF)
           ACMASS=AC*WN+DROP(AC, ZC)
   C
           CONSERVE MASS:
           DMASS=ACMASS-AFMASS
           TCN=((AP-DELTA A) *ELAB+(AT-B) *TA+DELTA A*TA)/AC
           TD = DELTA_A* (TCN+(3./5.)*EFERMI)+DELTA Z*Ec
   C
           CONSERVE TOTAL ENERGY:
           TF = (ACMASS+TCN*AC) - (DMASS+TD) - AFMASS
           PF = SQRT((AFMASS+TF)**2-AFMASS**2)
           dKE=ELAB-TF/AF
           dKE=SCALE*dKE
           IF(dKE.LT.O.) dKE=0.
           DP=PLAB-PF/AF
         END IF
   C
         Sigma in KE loss distribution:
          SIGMAKE = Temp*SQRT(9.*(AP-AFF)/AP)*(SQRT(.667*dKE/Temp)+1.)
         SIGMAKE=(SIGMAKE/ELAB)*100.
         RETURN
         END
₽₽ C
Œ
         FUNCTION HEAT (A0, AZ0, B0, BZ0, ELAB, A, AZ, B, BZ, C, CZ, TA, TB, TC, Temp)
E C
TI C
         CALCULATES THE ENERGY AND MOMENTUM OF THE THREE SOURCES A, B, C
LI C
ķ
         COMMON/RAPIDITY/YAO, YBO, YA, YB, YC
ų.
         DATA WN/931.504/
         DATA PI, EPS/3.14159,0.03/
  C
  С
         Transport parameters;
  C
                   Energy leaked to the spectators
            XΟ
  С
                   Longitudinal momentum degradation of spectators
            Y0
  C
            z_0
                   Tranveres momentum transfer:
  C
         DATA X0, Y0, Z0/.05, .25, 60./
  С
  C
         Sources A and Z numbers:
  С
         Note---These are estimated using Glauber theory. They are
  C
         impact-parameter averaged numbers!
  C
                   A is projectile spectator
  C
                   B is target spectator
  С
               and C is participant source.
  C
  C
         Masses of sources A, B, and C:
         A0MAS=A0*WN+DROP(A0,AZ0)
         BOMAS=B0*WN+DROP(B0,BZ0)
         AMAS=A*WN+DROP(A,AZ)
         BMAS=B*WN+DROP(B,BZ)
```

```
CMAS=C*WN+DROP(C,CZ)
   С
         EA=A0MAS+A0*ELAB
         PA=SQRT (EA**2-A0MAS**2)
         BETA0=PA/EA
         GAMMA0=1./SQRT(1.-BETA0**2)
         YA0=0.5*ALOG(ABS((1.+BETA0)/(1.-BETA0)))
         E0=EA+B0MAS
  C
  C
         In c.m. frame:
         V=PA/E0
         G=1./SQRT(1.-V**2)
         BETA CM=V
         GAMMA CM=G
         PA0=G* (PA-EA*V)
         E0=SQRT (E0**2-PA**2)
  C
  С
         Transport parameters X, Y, and Z, averaged over impact parameter:
  C
         [Recalculated 15 Dec. 1993]
  C
         T_FACTOR=(BETA_CM/BETA0)/GAMMA0**2
  C
         X = (1./2.) * X0 * T FACTOR
         Y = (1./2.) * Y0 * T FACTOR
         Z = (PI/4.) * Z0 * T_FACTOR
C
        Momenta of sources A, B, and C in the c.m. frame:
        PA = (1.-Y) *PAO*A/AO
        PB = -(1.-Y) * PAO * B/BO
        PC=-PA-PB
PAX = Z*A
PBX=-PAX
        PCX=-PAX-PBX
₽å C
TU C
        Iteration to find Q, the generated heat, conserving energy:
Ш
        N=1
Q=0.
v.
        DQ=0.
   10
        Q=Q+DQ
           N=N+1
           WA=AMAS+A*Q*X
           WB=BMAS+B*Q*X
           WC=CMAS+C*Q*(1.+(1.-X)*(A+B)/C)
           EA=SQRT (PAX**2+PA**2+WA**2)
           EB=SQRT (PBX**2+PB**2+WB**2)
           EC=SQRT (PCX**2+PC**2+WC**2)
           DQ=(E0-EA-EB-EC)/(A0+B0)
  C
        CHECK AVAILABLE ENERGY
           HEAT=DQ
           IF (Q.EQ.O..AND.DQ.LT.O.) RETURN
           IF (ABS(DQ).GT.EPS) GO TO 10
           E00=EA+EB+EC
  C
            IF ((E0-E00).GT.((A0+B0)*EPS)) PRINT 101, E0,E00
  C
  C
        Excitation Energy/nucleon:
        Temp=2./3.*Q*SQRT(T_FACTOR)
        HEAT=Q
           TA=(EA-AMAS)/A
           TB=(EB-BMAS)/B
```

罩

```
C
          Q=G*(EA+PA*V)
          PA=G* (PA+EA*V)
          EA=Q
   C
          Q=G* (EB+PB*V)
          PB=G* (PB+EB*V)
          EB=Q
   C
          Q=G*(EC+PC*V)
          PC=G* (PC+EC*V)
          EC=O
   C
         TA= (EA-AMAS) /A
         TB=(EB-BMAS)/B
         TC=(EC-CMAS)/C
         EE=EA+EB+EC
         P=PA+PB+PC
         E00=SQRT (EE**2-P**2)
   С
   С
          IF ((E0-E00).GT.((A0+B0)*EPS)) PRINT 101,E0,E00,EPS
   С
         RETURN
   С
        FORMAT (2X,20H ENERGY CONSERVATION,3F12.5)
    101
         FORMAT (/2X,6E12.4//2X,6E12.4)
    12
         END
         FUNCTION DROP (A, Z)
С
         CALCULATES THE LYSEKIL NUCLEAR MASS DEFECT+WIGNER+PAIRING+SHIFT
E C
=
         DATA A1,A2,C3,CAPPA/15.4941,17.9439,0.7053,1.7826/
-
         DATA C4/1.1533/
DATA WA, WN, WP/931.504,8.07169,7.28922/
         DATA WIG, D1, D2, SHIFT/30., 12., 10., 50./
222
         DROP=0.
ű
         IF (A.LT.0.9) RETURN
J<sub>LL</sub>
         A3=A**0.333333
         EN=(-A1*A+A2*A3**2)*(1.-CAPPA*(1.-2.*Z/A)**2)
         EC = (C3/A3 - C4/A) *Z**2
         W = (A-Z) *WN + Z*WP
         DROP=W
         IF (A.LE.4.) RETURN
         E=EN+EC+W
         T=ABS(1.-2.*Z/A)
        EW=WIG*T
         IA2=A/2.+0.1
        IF (A-2.*IA2-0.1) 10,10,20
     10 IZ2=Z/2.+0.1
        EP=D1/SQRT(A)-D2/A
         IF (Z-2.*IZ2-0.1) 11,11,15
     11 EW=EW-EP
        GO TO 30
     15 EW=EW+EP
        IF (IA2.EQ.2*IZ2) EW=EW+WIG/A
        GO TO 30
```

TC=(EC-CMAS)/C

```
20 EW=EW+D2/A
      30 CONTINUE
         E=E+EW+SHIFT/A
         DROP=E
   C
         RETURN
         END
   C
   C
         SUBROUTINE GLBR (AP, ZP, AT, ZT, AA, AAZ, BB, BBZ, CC, CCZ)
   C
   С
         Calculates (average) numbers of proj. and target participants according
         to Glauber theory, see, e.g., Tsao et al., PRC 47, 1257 (1993).
   C
   C
         INTEGER AP, ZP, AT, ZT, AA, AAZ, BB, BBZ, CC, CCZ
         INTEGER ZERO_A, ZERO_Z
   С
         DATA PI,RO/3.14159,1.36/
         DATA P13, P23/0.33333, 0.66667/
   C
         FACTOR1 = (AP**P13+AT**P13)**2
         FACTOR2 = AP**P23+2*AP**P13*AT**P13
   C
         Participants:
           AP P = AP * AT**P23 / FACTOR1
           AT P = AT * AP**P23 / FACTOR1
           ZPP = ZP * AT**P23 / FACTOR1
           ZT P = ZT * AP**P23 / FACTOR1
   С
         Participant source "C":
           CC = NINT(AP P) + NINT(AT P)
           CCZ = NINT(ZP_P) + NINT(ZT_P)
   С
   С
         Projectile spectator source "A":
           AA = AP - NINT(AP P)
           AAZ = ZP - NINT(ZP_P)
Li
  C
С
         Target spectator source "B":
           BB = AT - NINT(AT P)
           BBZ = ZT - NINT(ZT_P)
   С
   C
         Check baryon number conservation:
   C
         ZERO_A = (AP+AT) - (AA+BB+CC)
         ZERO_Z = (ZP+ZT) - (AAZ+BBZ+CCZ)
         IF (ZERO A.NE.O.OR.ZERO C.NE.O) THEN
   С
            PRINT*, ' ***Baryon Number Conservation***'
         END IF
   С
   C
           RETURN
```

\$

END

```
PROGRAM FLUX_DRIVER
```

Driver program for generating CREME96 model fluxes

IMPLICIT NONE

C C

С

INTEGER*4 IZMIN, IZMAX, IMODE, ITRANS, M

REAL*4 EMIN, EMAX, YEAR

CHARACTER*80 GTRANSFILE, TRAPDFILE, FLXFILE

INTEGER*4 MARR,NELM,VERSION_NUMBER,PROGRAM_CODE

PARAMETER (MARR=5000, NELM=92)

REAL*4 E, FLX

DIMENSION E (MARR), FLX (NELM, MARR)

CALL INIFLUX(IZMIN, IZMAX, EMIN, EMAX, YEAR, IMODE, ITRANS,

* GTRANSFILE, TRAPDFILE, FLXFILE)

CALL CREME96_FLUX(IZMIN, IZMAX, EMIN, EMAX, YEAR, IMODE, ITRANS,

* GTRANSFILE, TRAPDFILE,

* VERSION_NUMBER, PROGRAM CODE,

* M,E,FLX)

CALL OUTPUT_CREME96_FLUX(IZMIN, IZMAX, EMIN, EMAX,

* YEAR, IMODE, ITRANS,

* GTRANSFILE, TRAPDFILE,

* VERSION_NUMBER, PROGRAM CODE,

M, FLX, FLXFILE)

STOP

END

```
REAL FUNCTION GCR FLUX(IZIN, EEZ, YDUM, IPDUM)
            IMPLICIT NONE
            INTEGER I, IZIN, IPDUM, IMonthMean, Nmonths
   C
            Nmonths needs to be odd number to properly center on input month
            PARAMETER (Nmonths=1)
            REAL EEZ, YDUM, YEARAVG (Nmonths), GCR MONTHLY FLUX
   C
            Perform average over neighboring months to smooth out GCR fluxes
            GCR_FLUX=0.0
            IF (Nmonths .LE. 1) THEN
              YearAvg(1)=YDUM
              GCR_FLUX=GCR_MONTHLY_FLUX(IZIN, EEZ, YDUM, IPDUM)
              IF (Nmonths .LT. 1) THEN
                WRITE(*,222)
                FORMAT(1X,'Using number of months = 1 in GCR flux averages')
   222
ENDIF
           ELSE
Section Section
              IMonthMean=(Nmonths+1)/2
             DO I=1, NMonths
113
                YearAvg(I)=YDUM + FLOAT(I-IMonthMean)/12.0
há
                GCR FLUX=GCR_FLUX +
                          GCR MONTHLY FLUX(IZIN, EEZ, YearAvg(I), IPDUM)
<u>L</u>
             ENDDO
M. H.
GCR_FLUX=GCR_FLUX/FLOAT(Nmonths)
<u>Ē</u>
           ENDIF
           RETURN
           END
  С
  C
           REAL FUNCTION GCR MONTHLY FLUX(IZIN, EEZ, YDUM, IPDUM)
           IMPLICIT NONE
           INTEGER I, IZIN, IZ, IPDUM, IENT, K, J, Jyear
           INTEGER IYMIN, IYMAX, IY
           REAL*4 MO, EEZ, EN, R, RO, YDUM, PI
           REAL W(12,300), WF(3600), AN(92), d(28), b(28), to(28)
           REAL A(28), DD(28), ALPHA(28), GAMMA(28)
           REAL Z,T,T0,BETA,SINE,ARG1,ARG2,DELTA,F,DRDE,FLUX
           EQUIVALENCE (W(12,300), WF(3600))
  CX
             DIMENSION ENVAL(100), FVAL(100)
                                               !unused
```

For extrapolating wolf numbers to dates outside of data ranges,

```
REAL YearMin, YearMax
           REAL Tmin, SINE TERM, SGN, DT, Tmax, SUM, WS
           INTEGER MonthMin, MonthMax, IBmin, IBmax, IB, N, STAT, CREME96_OPEN
   C
           made separate array for reading wolf number file. This allows
   C
           input wolf numbers to start and end and any particular date,
   C
           and preserves the original structure with W and WF variables.
           REAL WOLFTMP (3600)
   C
   C
           ENTER Tmax AND THE REST MASS OF A NUCLEON, MO
   C
           DATA Tmax/14.5/,M0/931.162/
   C
   C
           ATOMIC MASS TABLE
   С
           DATA (AN(I), I=1, 92)/1., 4., 6.9,
        1 9.,10.8,12.,14.,16.,19.,20.2,23.,24.3,27.,28.,
        2 31.,32.,35.5,39.9,39.,40.,45.,47.9,50.9,
        3 52.,54.9,55.8,58.9,58.7,63.5,65.4,69.7,72.6,
        4 74.9,79.,79.9,83.8,85.5,87.6,88.9,91.2,92.9,95.9,97.,101.,
        5 102.9,106.4,107.9,112.4,114.8,118.7,121.8,127.6,126.9,131.3,
        6 132.9,137.3,138.9,140.1,140.9,144.2,145.,150.4,152.,157.3,
        7 158.9,162.5,164.9,167.3,168.9,173.,175.,178.5,180.9,183.9,
        8 186.2,190.2,192.2,195.1,197.,200.6,204.4,207.2,209.,209.,
        9 210.,222.,223.,226.,227.,232.,231.,238./
T
  С
С
           TABLE 1 FROM NYMMIK ET AL.
C
14
           DATA b/28*1.2/,to/28*1982.5/
  C
TOTAL
TOTAL
⊨ C
           TABLE 2 FROM NYMMIK ET AL. NOTE: D IN THE TABLE IS DD HERE
TU C
IJ
           data A/1.,4.,6.9,9.,10.8,12.,14.,16.,19.,20.2,23.,24.3,27.,
        * 28.1,31.,32.1,35.4,39.9,39.1,40.1,44.9,47.9,50.9,52.,54.9,
* 55.8,58.9,58.7/
Q
          DATA DD/2.0E04,3.5E03,1.7E01,1.6E01,5.1E01,9.6E01,3.5E01,
* 8.4E01,3.6E00,1.5E01,4.2E00,1.8E01,3.9E00,1.2E01,1.0E00,
        * 2.7E00,1.2E00,2.3E00,1.8E00,2.6E00,6.9E-01,2.5E00,1.13E00,
       * 2.1E00,1.04E00,9.2E00,8.7E-02,4.5E-01/
          DATA ALPHA/3.,3.,3.4,4.5,3.9,3.1,3.6,3.0,3.8,3.1,3.4,3.0,
       * 3.2,3.0,4.0,3.4,4.5,4.5,4.2,3.2,3.6,3.6,3.3,3.3,3.3,3.0,3.1,
       * 4.0,3.2/
          DATA GAMMA/2.75,2.75,2.70,2.90,3.00,2.75,2.90,2.70,3.00,
       * 2.75,2.90,2.70,2.80,2.65,2.95,2.70,3.00,2.90,3.00,2.75,
       * 2.90,2.95,2.90,2.85,2.70,2.60,2.75,2.60/
  С
  C-----
  C
  С
          Routine extended to Z >28 using relative abundances from CREME,
  С
          applied to the Fe spectrum calculated here:
  С
          REAL RCREME (92)
  C
          THE ELEMENTAL RATIOS HAVE BEEN EXTENDED TO URANIUM USING
  С
          THE HEAO-3 DATA INTERPRETED WITH CAMERON'S ABUNDANCES AND
  C
          THE RESULTS OF COSMIC RAY PROPAGATION CALCULATIONS
```

assuming a 22 year periodicity. PRB.

```
A.G.W.CAMERON, HARVARD-SMITHSONIAN CENTER FOR ASTROPHYSICS
   С
   С
            PREPRINT SERIES NO. 1357, 1980, AND TSAO ET AL., PROC. OF
   C
           THE 17TH INTL. COSMIC RAY CONF., VOL. 9, P130-33, PARIS, 1981).
   C
           TABLE 7 IN CREME REPORT.
   С
   C
           Revised 6/19/92 by AJT, using latest combined HEAO-3/Ariel
   C
           abundances, as reported by Binns et al. Ap.J 346,997-1009, 1989.
   C
           NOTE: since these measurements cannot always resolve individual
   C
                   elements, the numbers here preserve the even-odd and
                   intra-group ratios from CREME IV.
   C
   C
           DATA (RCREME(I), I=29, 92)/
        1 6.8E-4,8.8E-4,6.5E-5,1.4E-4,8.9E-6,5.2E-5,
        1 9.7e-6,2.7E-5,8.8E-6,2.9E-5,6.5E-6,1.6E-5,2.9E-6,8.1E-6,9.5E-7,
        2 3.1e-6,1.6E-6,4.6E-6,1.5E-6,4.0E-6,8.8E-7,4.7E-6,9.9E-7,5.7E-6,
        3 1.1e-6,2.7e-6,6.5E-7,6.7E-6,6.0E-7,1.8E-6,4.3E-7,1.6E-6,1.9E-7,
        4 1.8e-6,3.1e-7,1.4E-6,3.5E-7,1.4E-6,5.3E-7,8.8E-7,1.8E-7,8.9E-7,
        5 1.3e-7,8.1e-7,7.3E-8,8.1E-7,2.8E-7,1.2E-6,7.9E-7,1.5E-6,2.8E-7,
        6 4.9e-7,1.5e-7,1.4E-6,7.3E-8,0.,0.,0.,0.,0.,8.1E-8,0.,4.9E-8/
   C
           IZ=IZIN
           IF (IZ.GT.28) IZ=26
Hard Hard Lines
           IF (IENT.EQ.0) THEN
           IENT=1
d(1) = 0.012
do i=2,28
į.
           d(i) = d(1) *an(i) /float(i)
           end do
ì.i.
TI C
           CALCULATE PI
LU C
i.e.
           PI=4.0*ATAN(1.0)
i C
≒,j C
           OPEN FILE OF MONTHLY AVERAGE WOLF NUMBERS FROM MCKINNON AT NOAA
   С
           AND READ THEM IN
  C
          use file starting in 1950
  C
  C
            OPEN (UNIT=60, READONLY, SHARED, STATUS='OLD',
  C
                 FILE='CREME96:WOLF.DAT')
           stat = creme96_open('wolf.dat','cr96tables',60,'old')
           IF (ydum .LT. 1950) THEN
             WRITE(*,111)
  111
             FORMAT(1X,'Warning, GCR results are unreliable before 1950')
           ENDIF
          Modified read, where K is set to maximum array size. At present,
  С
           this is 3600, allowing specification of 300 years.
          DO K = 1,3600
```

(BINNS ET AL., AP. J., VOL. 247, L115-L118,1981,

C

200

```
THE REAL PROPERTY.
įmi
E C
O
```

```
С
              For determining bounds of Wolf number file. If entered date is
   C
              outside of these bounds, the wolf numbers are extrapolated
   C
              assuming a 22 year periodicity.
              MonthMax=I
              YearMax=FLOAT(Jyear) + FLOAT(MonthMax-1)/12.0
              READ (60, 1, END=2) JYEAR, I, WOLFTMP (K)
              J = (JYEAR - 1749) + 1
              W(I,J) = WOLFTMP(K)
              IF (K .EQ. 1) THEN
                MonthMin=I
                YearMin=FLOAT(Jyear) + FLOAT(MonthMin-1)/12.0
                MonthMax=I
                YearMax=FLOAT(Jyear) + FLOAT(MonthMax-1)/12.0
              ENDIF
           ENDDO
   1
           FORMAT (14, 1X, 12, F6.1)
           CONTINUE
           CLOSE (60)
           ENDIF
                      !IENT = 0 OPTION
           COMPUTE THE BOUNDS OF THE WOLF NUMBER ARRAY. This update
   C
           is to be used in smoothing algorithm, so that check if out of
           bounds. Also used by CheckDates routine, in order to handle
   C
  C
           transition in wolf number array at the center of each month,
TU C
           e.g. wolf number changes from month 6 (June) to month 7 (July)
LJ C
           on the 15 June, as used in IB index.
           IYMIN=INT(YearMin)
           IBMIN= (IYMIN-1749) *12+12.* (YearMin-IYMIN) +.5+1
           IYMAX=INT(YearMax)
           IBMAX=(IYMAX-1749)*12+12.*(YearMax-IYMAX)+.5+1
           Z=IZ
           GCR_MONTHLY_FLUX=0.0
   C
   C
           CALCULATE DT (MONTH) FOR YEAR T
   C
           T=YDUM
                                YDUM will now be used to pass the year of
           IY=INT(T)
                            1
                                interest to this function. R. Witt 6/22/95
   C
   C
           COMPUTE THE LOCATION, IB, OF THE WOLF NUMBER FOR TIME T IN
   C
           THE W ARRAY
   С
           IB = (IY-1749) *12+12.*(T-IY) +.5+1
```

```
Routine checks if T is within Wolf number bounds. If not, adjusts
  C
          T and IB assuming a periodicity of 22 years.
  C
          Note that period which crosses beginning year in wolf number data
          boundary has been handled as a special case in the DO 70 and DO 71
  С
  С
          loops below.
          CAll CheckDate(T, YearMin, Yearmax, IB, IBmin, IBmax)
  C
          VERSION OF TO IN THE PAPER
  С
          T0=1978.5
          IF(T.GE.1985.) T0=1976
  C
          EN=EEZ
  С
  С
          CONVERT EN TO RIGIDITY IN GV, AS R
  C
          R = (AN(IZ)/Z) * (EN*EN+2.*M0*EN) **.5/1000.
  С
  C
          CALCULATE DT (MONTH) FOR R IN YEAR T
  C
          Tmin = 5.3/R**0.3
          SINE TERM=sin(2.*PI*(t-t0)/22.)
          IF (SINE TERM.GE.O.) SGN=1.
          IF(SINE_TERM.LT.0.) SGN=-1.
DT = (Tmax + Tmin) / 2. + ((Tmax - Tmin) / 2.) * SGN*
       1 (ABS(SINE_TERM)) ** (1./3.)
          CALCULATE THE SMOOTHED WOLF NUMBER, WS
          N=DT+0.5
SUM=0.
i.i.
          DO 70 K=1, N
IF ((IB-K) .GE. IBMIN) SUM=SUM+K*WF(IB-K)
LI C
            Extrapolate backwards using 22 year solar cycle pattern
IF ((IB-K) .LT. IBMIN) SUM=SUM+K*
WF(IB-K+12*22)
70
          CONTINUE
          DO 71 K=N+1,2*N-1
            IF ((IB-K) .GE. IBMIN) SUM=SUM+(2*N-K)*WF(IB-K)
  C
            Extrapolate backwards using 22 year repeating pattern
            IF ((IB-K) .LT. IBMIN) SUM=SUM+(2*N-K)*
                WF(IB-K+12*22)
  71
          CONTINUE
          WS=SUM/(N*N)
  C
  C
          COMPUTE THE MODULATION POTENTIAL
  C
          R0=0.375+3E-4*WS**1.445
 C
 C
          COMPUTE BETA
 C
          BETA=SQRT(1-(EN/M0+1)**(-2))
 C
 C
          COMMPUTE DELTA
 C
```

C

25

```
SINE=SIN(2.*PI*(T-TO(IZ))/22.)
           IF(SINE.GE.O.) SGN=1.0
           IF(SINE.LT.0.) SGN=-1.0
           SINE=ABS(SINE)
           Inserted to avoid floating underflows.
   С
           ARG1 = -BETA * R/D(IZ)
           ARG2 = - BETA*R/R0
           IF (ABS(ARG1) .LE. 1.0E-20) ARG1=0.0
           IF (ABS(ARG2) .LE. 1.0E-20) ARG2=0.0
           DELTA=5.5*ABS(1-B(IZ)*EXP(ARG1))+
          (1.13*BETA*R/R0)*(SGN*SINE**(1./3.))*EXP(ARG2)
           IF (ABS(DELTA) .LE. 1.0E-20) DELTA=0.0 !also to avoid underflows
   C
           COMPUTE THE FLUX
   C
   C
           F = (DD(IZ) *BETA * *ALPHA(IZ)) / R * *GAMMA(IZ)
           F=F*(R/(R+R0))**DELTA
   C
   C
           COMPUTE dR/dE
  C
           dRdE = (AN(IZ) / (Z*1000.)) * (EN+M0) / ((EN*EN+2.*M0*EN) **.5)
С
           CONVERT FROM PER GV TO PER MeV/nuc
           FLUX=F*dRdE
           GCR_MONTHLY_FLUX=GCR_MONTHLY_FLUX+FLUX
Ш
<u>___</u>
22
           Scale relative to Fe for Z > 28:
ğ-zşk
           IF (IZIN.GT.28) GCR MONTHLY FLUX=GCR MONTHLY FLUX*RCREME(IZIN)
T.
           RETURN
L
           END
7,1
           SUBROUTINE CheckDate (Year, YearMin, Yearmax, IB, IBmin, IBmax)
           Routine checks if T is within Wolf number bounds. If not, adjusts
   C
           T to being in bounds assuming a 22.0 year periodicity.
   C
   С
           YearMin & YearMax are unused in this routine, but are included
           as arguments for possible future use.
           IMPLICIT NONE
           REAL Year, YearMin, Yearmax
           INTEGER Ncycle, IBmin, IBmax, IB, IBlow, IBhigh, IBnew
   C
           fix wolf number range for extrapolating to be July 1970 to JUNE 1992
           this algorithm assumes a 22 year periodicity.
   C
           DATA IBlow, IBhigh/2659, 2922/
   C------
```

IF ((IB .GE. IBmin) .AND. (IB .LE. IBmax)) THEN

```
Dates are o.k., don't need to adjust YEAR or IB
RETURN

ELSEIF (IB .GT. IBmax) THEN
IBnew=IBlow+MOD(IB-IBhigh-1,IBhigh-IBlow+1)
Ncycle=(IB-IBhigh-1)/(IBhigh-IBlow+1)+1
Year=Year-Ncycle*22.0
IB=IBnew
ELSEIF (IB .LT. IBmin) THEN
IBnew=IBhigh-MOD(IBlow-IB-1,IBhigh-IBlow+1)
Ncycle=(IBlow-IB-1)/(IBhigh-IBlow+1)+1
Year=Year+Ncycle*22.0
IB=IBnew
ENDIF

RETURN
END
```

```
SUBROUTINE Geomag96 (OrbIncl, Apogee, Perigee, AscNodeLong,
      #
                       AscNodeDisp, PerigDisp, Zenith, Azimuth, UTtimeInit,
      #
                        Stormy, Shadow, PreCalcGTFs, IPreCalc,
      #
                       RigBins, TransFunc, Year, XLbounds, ILbins)
         IMPLICIT NONE
         INTEGER J, Jmax, L, Ndays, NorbSteps, IPreCalc, Nrigs, NLvals
         PARAMETER (Ndays=7, NorbSteps=200, Nrigs=1001, NLvals=10)
С
        Now REAL to properly handle omnidirectional averaging.
C
        Only the Earth's geometric shadow is included in the generic
C
        omnidirectional averaging at present.
        REAL MAT (Nrigs, NLvals), TransInc
        INTEGER IDEX
        DATA TransInc/1.0/
        REAL RigBins (Nrigs), TransFunc (Nrigs, NLvals)
        LOGICAL Shadow, Stormy, PreCalcGTFs, GridInit, INIGRID
С
        Initial Orbital & lookout direction input parameters set
        in GTFDriverInput
        REAL OrbIncl, Apogee, Perigee, AscNodeLong, AscNodeDisp, PerigDisp
        REAL Zenith, Azimuth, C, Cgrid, Csupress, DeltaNymmik
        REAL UTtimeInit, UTtime, TimeLocal
        REAL Time, Period, Step
        Parameters along each orbital step
        REAL Zlat, Zlon, Alt
        INTEGER ILbins, ILbin, ICODE, NperLbin(NLvals)
        REAL Year, XLval, BBO, XLbounds (NLvals), XLinfinite
        PARAMETER (XLinfinite=1.0E+06)
        REAL Grid80Lval, RatioL
        LOGICAL UseLapprox
        Initializations
C
        DO L=1, NL vals
         NperLbin(L)=0
         DO J=1, Nrigs
           MAT(J,L)=0.0
         ENDDO
        ENDDO
        TransInc=1.0
        Choice of original geomagnetic storm option or pre-calculated GTF
C
C
        option. These are mutually exclusive now. Note that "Stormy" applies
C
        to updated Grid, and thus will be applied on top of the Nymmik
```

```
IF (PreCalcGTFs) THEN
                     The pre-calculated GTFs have not been divided into L-bins.
    C
                     This may be a useful option to include in future updates.
              CALL GetPreCalcGTF(IPreCalc,RigBins,TransFunc)
              RETURN !could just use subsequent RETURN, since this IF statement
                      !skips all lines before the subsequent RETURN
            ELSE !calculate GTF if not using pre-calculated ones
              Initialize Orbit routine
    C
              CALL Orbit(1, Period, ZLon, ZLat, Alt, Apogee, Perigee, OrbIncl,
                           AscNodeLong, AscNodeDisp, PerigDisp)
         #
    C
              Initialize cutoff grid.
    C
    C
              IF (.NOT. GridInit) GridInit=INIGRID
              Compute the total number of steps in "Ndays" days if we make
   C
"Norbsteps" steps per orbit. Use 2 days and 200 steps per orbit
    C
              presently.
    С
    С
              JMAX=INT(Ndays* NorbSteps*86400./PERIOD + 1.5)
              Compute the step size in seconds.
    C
              STEP=PERIOD/FLOAT (NorbSteps)
    С
              Compute the vertical cutoff at the spacecraft
              position for every time step.
             DO J=1, JMAX
              time=FLOAT(j-1)*step
              CALL Orbit(2, Time, ZLon, ZLat, Alt, Apogee, Perigee, OrbIncl,
                            AscNodeLong, AscNodeDisp, PerigDisp)
            Now calculate geomagnetic cutoff from the Grid. Perform before
            L-value calculation, and see similarity
    C
              CALL GET CUTOFF (ZLAT, ZLON, ALT, Azimuth, Zenith, C)
              Cgrid=C
              IF ( XLbounds(2) .LT. XLinfinite .OR.
                    ( XLbounds(2) .GE. XLinfinite .AND.
         &
                     XLbounds(1) .GT. 0.0 ) ) THEN
                CALL GridApproxLval (Cgrid, XLbounds, ILbins, Grid80Lval,
                          UseLapprox)
         æ
```

IF (UseLapprox) THEN
 XLval=Grid80Lval

correction for high inclination orbits if that option is chosen.

C

ELSE

```
CALL GET BLCOORDS (Year, Zlat, Zlon, Alt, XLval, BB0, ICODE)
            ENDIF
            IF (XLval .GT. 99999.0) XLval=99999.0
            CALL GetLbin (XLval, XLbounds, ILbins, ILbin)
            IF (ILbin .GE. 1 .AND. ILbin .LE. NLvals)
                NperLbin(ILbin) = NperLbin(ILbin) +1
     &
          ELSE
C
            If no L-bins are specified or 1 L-bin is specified
С
            and the lower bound is L = 0, use only the first
            element of the array. In this case, the following
C
C
            sum should equal JMAX once the stepping through the
C
            orbit is completed.
            ILbin=1
            NperLbin(Ilbin) = NperLbin(Ilbin) +1
          ENDIF
          CALL ConvertTime (time, UTtimeInit, UTtime, Zlon, Period,
                           TimeLocal)
                      IF (Cgrid .GT. 0.0) THEN
          CALL Nymmik (C, TimeLocal, DeltaNymmik)
          C=C/(1+DeltaNymmik)
          IF (Stormy) THEN
C
              Now apply cutoff suppression during large magnetic storms,
С
              as described by Adams, et al. (1981).
            Csupress = .54*EXP(-Cgrid/2.9)
            C=C-Csupress
            IF (C .LT. 0.) C=0.0 !lowest cutoffs are defined to be 0
          ENDIF
                            !applying Stormy correction
         ENDIF
                           !checking that Cgrid (grid cutoff) > 0.0
\mathsf{C}
C
        Histogram cutoffs in 0.02 GV steps. Since only allow vertical and
C
        western cutoffs, all IDEX should be in bounds, since C vert < 20 GV.
C
        Note that the transmission function is an integral spectra of
C
        cutoffs < Rigidity. See CALCULATE TRANS FUNC for algorithm
C
        which assigns rigidities for bins.
         IF (C .EQ. 0.0) THEN
           IDEX=1
           IDEX=INT(C*50.)+2
         ENDIF
```

This is a correction for the earth's shadow on the spacecraft

```
according to simple geometrical optics. Have made MAT real, and
С
        apply to each point in calculation, in order to handle correction
C
        properly for non-circular orbits. This routine has been designed
C
        to always apply the Earth's shadow, although the technique will likely
C
        be modified before 1997.
C
C
         IF (Shadow) THEN
            TransInc=(1.-0.5*(1.-((6371.2+ALT)**2.
                             -(6371.2)**2.)**.5/(6371.2+ALT)))
     1
         ENDIF !applying Earth's shadow correction
         IF (ILbin .GE. 1 .AND. ILbin .LE. NLvals)
             MAT(IDEX, ILbin) = MAT(IDEX, ILbin) + TransInc
                  !for number of orbital steps
         ENDDO
C
        Now calculate transmission function.
C
C
         CALL CALCULATE TRANS_FUNC(Jmax, MAT, RigBins, NperLbin, TransFunc)
                !for using either pre-calculated GTF or GRID-based options
        ENDIF
        RETURN
        END
        SUBROUTINE GET CUTOFF (ZLAT, ZLON, ALT, AZ, ZE, C)
C
C
        For input ZLAT, ZLON, ALT, AZ, ZE, calculates cutoff C (in GV)
            ZLON = geocentric longitude of spacecraft position (deg)
С
С
            ZLAT = geocentric latitude of spacecraft position (deg)
            ALT = spacecraft altitude (km)
C
                 = azimuth of particle wrt spacecraft (deg)
C
            AZ
                 = zenith angle of particle wrt spacecraft (deg)
С
            zE
C
C
        Routine modified 3/5/90:
C
          In JHA's original version of this coding, he first calculated the
C
             vertical cutoff at the 4 grid corners at 20 km altitude,
C
             re-scaled via Stormer theory to orbital altitude and orientation,
C
C
             and then averaged the four. This procedure involved 5 calls
             to subroutine FUNCTION STORMER.
С
С
C
          In the modified coding, the 20 km vertical cutoffs are first averaged,
             and then re-scaled via Stormer theory to orbital altitude and
С
             orientation. This procedure involves only 2 calls to STORMER.
С
C
             also gives a smoother transmission function.
C
        12/16/92, Fixed XORB bug, so that XORB is calculated for
С
C
             all latitudes. RGRD is now a parameter.
C
        12/20/95, set cutoffs that are negative to 0.0.
        IMPLICIT NONE
        REAL ZLAT, ZLON, ALT, AZ, ZE, C
```

REAL AZG, ZEG, RGRD

```
DATA AZG/0./, ZEG/0./
        PARAMETER (RGRD=1.0031392126) !equivalent to 6391.2/6371.2
        REAL CUTOFF (33,72), CN, CS
        COMMON/CUTOFF80/CUTOFF, CN, CS
        INTEGER ILO, IUP, JLO, JUP
        REAL ZI, ZJ, XORB, DI, DJ, SC, SG
        REAL Y1, Y2, Y3, Y4, CL
        REAL STORMER
C-----
        COMPUTE THE TABULAR POSITION OF THE VERTICAL CUTOFF.
С
C
        ZI=ZLAT/5.+17.
        ZJ=ZLON/5.+1.
        ILO=INT(ZI)
        IUP=ILO+1
        JLO=INT(ZJ)
        IF(JLO.EQ.73) JLO=1
        JUP=JLO+1
        IF (JUP.EQ.73) JUP=1
С
        INTERPOLATE THE VERTICAL CUTOFF TO THE EXACT LOCATION
C
        OF THE SPACECRAFT USING STORMER THEORY.
C
C
        XORB = (6371.2 + ALT) / 6371.2
        IF (ABS (ZLAT) .GE.80.) GO TO 100
        DI=ZI-FLOAT(ILO)
        DJ=ZJ-FLOAT (JLO)
        SC=STORMER (ZLAT, ZLON, XORB, AZ, ZE)
        SG=STORMER (ZLAT, ZLON, RGRD, AZG, ZEG)
C
        Vertical cutoffs
        Y1=CUTOFF(ILO, JLO)
        Y2=CUTOFF(IUP,JLO)
        Y3=CUTOFF(ILO, JUP)
         Y4=CUTOFF(IUP, JUP)
         C = (1.-DI) * (1.-DJ) * Y1 + (1.-DI) * DJ * Y3 + DI * (1.-DJ) * Y2 + DI * DJ * Y4
        C=SC*C/SG
        GO TO 200
C
C
         FOR ABS(LATITUDE).GT.80 USE THE CUTOFFS AT THE POLE INSTEAD OF
С
         THE CUTOFFS AT FOUR NEARBY LOCATIONS.
С
 100
         CONTINUE
         DJ=ZJ-FLOAT (JLO)
         SC=STORMER(ZLAT, ZLON, XORB, AZ, ZE)
         SG=STORMER (ZLAT, ZLON, RGRD, AZG, ZEG)
         IF(ZLAT.LE.-80.)GO TO 110
         DI=ZI-33.
```

CL=DJ*CUTOFF(33,JUP)+(1.-DJ)*CUTOFF(33,JLO)

```
C = (DI*CN+(2.-DI)*CL)/2.
       C=SC*C/SG
       GO TO 200
110
       CONTINUE
       DI=1.-ZI
       CL=DJ*CUTOFF(1,JUP)+(1.-DJ)*CUTOFF(1,JLO)
       C = (DI*CS+(2.-DI)*CL)/2.
       C=SC*C/SG
200
       CONTINUE
       IF (C .LT. 0.) C=0.0 !added 12-20-95
       RETURN
       END
C-----
       SUBROUTINE GetLbin(XLval, XLbounds, ILbins, ILbin)
       IMPLICIT NONE
       INTEGER ILbins, ILbin, NLvals, L
       PARAMETER (NLvals=10)
       REAL XLval, XLbounds (NLvals)
       LOGICAL FindLbin
       No attempt is made to eliminate "unphysical" or "approximate"
C
       L-values using the ICODE returned from GET_BLCOORDS, since any
C
       analyses using L-values are likely to handle these locations
C
        "as is", i.e. with the calculated L-value.
C
        FindLbin=.TRUE.
        ILbin=0
        DO L=1, ILbins
         IF (FindLbin) THEN
           IF (L .LT. NLvals) THEN
             IF ( (XLval .GE. XLbounds(L)) .AND.
             (XLval .LT. XLbounds(L+1)) ) THEN
               ILbin=L
               FindLbin=.FALSE.
             ENDIF
           ELSE !special handling of L=NLvals case
             IF (XLval .GE. XLbounds(L))THEN
               ILbin=L
               FindLbin=.FALSE.
             ENDIF
           ENDIF !checking of each L-bin
                  !for FINDLbin logical
          ENDIF
         ENDDO
         RETURN
         END
```

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C C C

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C

C

C My attempt to make this into a more modern FUNCTION, including the use of IMPLICIT NONE. 5-7-96, PRB.

REAL FUNCTION STORMER (GCLATD, GCLOND, RGC, AZ, ZE)

WE DID NOT WRITE THIS SUBROUTINE. WE HAVE MADE NO CHANGES IN IT IN 1984.

May 1996 comments and status, PRB.

- 1. Note that this FUNCTION uses the 1975 IGRF field plus drifts. In principle, the 1980 IGRF/DGRF coefficients would be more appropriate, since the STORMER corrections are applied to the 1980 grid. In future years, we intend to replace the GRID results with a 1990 grid, and will modify this routine accordingly.
- 2. The coefficients are also listed in inverted order, compared with more recent tabulations, e.g. G01 is generally listed as G10 in more recent tabulations.
- 3. This routine HAS NOT been converted to IMPLICIT NONE, due to the historical nature of the coding.

IMPLICIT NONE

REAL RED, EDLAT, AZM, ZEM, GAMMA

COMMON/KARL/RED, EDLAT, AZM, ZEM, GAMMA

Need to determine usage of DPEC(U), 5-7-96, PRB. REAL DPEC,U,ZEDRTL,ZRTL

INTEGER JDATA, NOPT

REAL PI, RAD, PIO2, TWOPI, SQRT3, DT

REAL G01, G02, G11, G12, H11, H12, G22, H22, H0, H0SQ

REAL ELO, EL1, EL2, E, XEDFGC, YEDFGC, ZEDFGC, REDFGC

REAL THETA, THETAD, PHI, PHID, CP, SP, ST, CT, CPCT, CPST, SPCT, SPST

REAL RIER, RIKM, ERAD, THIRAD, THIDEG, PHIRAD, PHIDEG

REAL XGMED, YGMED, ZGMED, ZDEDNP, RGC, XODNP, YODNP, ZODNP, DODNP

REAL DIFLA, PHINOF, TNOF, SGCLATD, CGCLATD, GCLATD, SGCLOND, GCLOND

REAL CGCLOND, XGC, YGC, ZGC, GCT, SGCT, CGCT, GCROT, SGCROT, CGCROT

REAL XRL, YRL, ZRL, XEDRL, YEDRL, ZEDRL, XEDRTL, YEDRTL, XRTL, YRTL

REAL XEDP, YEDP, ZEDP, XODNPR, YODNPR, ZODNPR, XODNPT, YODNPT, ZODNPT

REAL ROTM, SROTM, CROTM, ROTMD, PLAZ, AZ, TLZE, ZE, SPLAZ, CPLAZ

REAL STLZE, CTLZE, XLD, YLD, ZLD, CA, A, SA, ADEG, XLP, YLP, ZLP, CB, B, SB

REAL BDEG, XLPP, YLPP, ZLPP, ZLDM, XLDM, YLDM, SMALL, PAZM

REAL XED1, YED1, ZED1, ZED2, COSLDA

THIS FUNCTION TRANSFORMS A GEOGRAPHIC LOCATION AND ARRIVAL DIRECTION INTO OFFSET DIPOLE COORDINATES, THEN COMPUTES THE STORMER CUTOFF IN GV AND RETURNS THE RESULT. THE OFFSET DIPOLE COORDINATES ARE AVAILABLE IN THE COMMON BLOCK /KARL/.

00000

```
GCLATD IS GEOCENTRIC LATITUDE IN DEGREES
C
           GCLONG IS GEOCENTRIC LONGITUDE IN DEGREES
C
                  IS RADIAL DISTANCE FROM GEOCENTER IN EARTH RADII
           RGC
C
                  IS GEOGRAPHIC AZIMUTH
С
           AZ
           ZE
                  IS GEOGRAPHIC ZENITH
C
                  IS RADIAL DISTANCE FROM OFFSET DIPOLE POSITION IN
           RED
C
                                            EARTH RADII
C
        EDLAT IS THE GEOMAGNETIC LATITUDE IN OFFSET DIPOLE COORDINATES
C
                  IS GEOMAGNETIC AZIMUTH IN OFFSET DIPOLE COORDINATES
С
                  IS GEOMAGNETIC ZENITH IN OFFSET DIPOLE COORDINATES
C
           ZEM
           GAMMA IS GAMMA ANGLE MEASURED FROM MAGNETIC EAST
С
C
       DATA JDATA, NOPT/2*0/, PI, RAD, PIO2, XEDFGC, YEDFGC, ZEDFGC, CP, SP
     1 ,ST,CPCT,CPST,SPCT,SPST,XGMED,YGMED,ZGMED/16*-8000./
       DATA SMALL/1.0E-35/
      In declaration section now, 5-96, PRB.
С
      DATA ERAD, THETAD, PHID, R1KM, TH1DEG, PH1DEG/6371.2,
     1 11.4354,-290.2392,450.2586,72.8278,148.7753/
       IF (JDATA.EQ.77)
                         GO TO 10
         PI = ACOS(-1.0)
         RAD = 180.0/PI
         PIO2 = PI/2.0
         TWOPI = PI*2.0
       NOPT = 0
       SQRT3 = SQRT(3.0)
           ENTER GEOMAGNETIC DATA,
                                      IGRF 1975
C
C
             SEE JGR, 81, 5163, 1976
                                      DT IS NUMBER OF YEARS SINCE 1975
С
       DT = 5.0
       G01 = -30186.0 + 25.6*DT
       G02 = -1898.0 - 24.9*DT
       G11 = -2036.0 + 10.0*DT
       G12 = 2997.0 +
                         0.7*DT
              5735.0 - 10.2*DT
       H11 =
       H12 = -2124.0 -
                           3.0*DT
       G22 = 1551.0 + 4.3*DT
       H22 =
                -37.0 - 18.9*DT
       IF(NOPT.EQ.1) PRINT 1000, G01, G02, G11, G12, G22, H11, H12, H22
         COMPUTE POSITION OF OFFSET DIPOLE
C
       H0 = SQRT(G01*G01+G11*G11+H11*H11)
       HOSQ = HO*HO
       EL0 = 2.0*G01*G02+(G11*G12+H11*H12)*SQRT3
       EL1 = -G11*G02+(G01*G12+G11*G22+H11*H22)*SQRT3
       EL2 = -H11*G02+(G01*H12-H11*G22+G11*H22)*SQRT3
       E = (EL0*G01+EL1*G11+EL2*H11)*4.0*H0SQ
C
       E = (EL0*G01+EL1*G11+EL2*H11)/(4.0*H0SQ)
       IF (NOPT.EQ.1) PRINT 1011, EL0, EL1, EL2, E, H0
 1011 FORMAT (1H , 8E15.5)
        XEDFGC = ERAD*(EL1-G11*E)/(3.0*HOSQ)
C
                    (EL1-G11*E)/(3.0*H0SQ)
       XEDFGC =
C
        YEDFGC = ERAD*(EL2-H11*E)/(3.0*H0SQ)
                     (EL2-H11*E)/(3.0*H0SQ)
C
        ZEDFGC = ERAD*(EL0-G01*E)/(3.0*H0SQ)
                     (ELO-GO1*E)/(3.0*HOSQ)
       REDFGC = SQRT (XEDFGC*XEDFGC+YEDFGC*YEDFGC+ZEDFGC)
```

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IF (NOPT.EQ.1) PRINT 3001, XEDFGC, YEDFGC, ZEDFGC, REDFGC
 3001 FORMAT (1H , 4F10.4,
                                  3X, 'XEDFGC, YEDFGC, ZEDFGC, REDFGC')
 1000 FORMAT (1H , 10F13.5)
 1010 FORMAT(1H0, 8F15.5/1H, 8F15.5)
       THETA = THETAD/RAD
       PHI = PHID/RAD
       CP = COS(PHI)
       SP = SIN(PHI)
       ST = SIN(THETA)
       CT = COS(THETA)
       CPCT = CP*CT
       CPST = CP*ST
       SPCT = SP*CT
       SPST = SP*ST
       R1ER = R1KM/ERAD
       TH1RAD = TH1DEG/RAD
       PH1RAD = PH1DEG/RAD
       IF (NOPT.EQ.1) PRINT 1000, R1KM, TH1DEG, PH1DEG, R1ER, TH1RAD, PH1RAD
       XGMED = XEDFGC*CPCT -YEDFGC*SPCT -ZEDFGC*ST
       YGMED = XEDFGC*SP +YEDFGC*CP
       ZGMED = XEDFGC*CPST -YEDFGC*SPST +ZEDFGC*CT
       IF (NOPT.EQ.1) PRINT 3002, XGMED, YGMED, ZGMED
 3002 FORMAT(1H , 3F10.4, 13X, 'XGMED, YGMED, ZGMED')
       IF(NOPT.EQ.1) PRINT 1010, CP, SP, CT, ST, CPCT, CPST, SPCT, SPST
       JDATA = 77
   10 CONTINUE
          ITERATE TO FIND COORDINATES OF OFFSET NORTH DIPOLE AT ANY
C
C
       FIRST GUESS FIND OFFSET NORTH DIPOLE AT DISTANCE RGC
       ZDEDNP = RGC
  100 XODNP = XGMED*CPCT + YGMED*SP + ZDEDNP*CPST
       YODNP = -XGMED*SPCT + YGMED*CP - ZDEDNP*SPST
                                      + ZDEDNP*CT
       ZODNP = -XGMED*SP
       DODNP = SQRT (XODNP*XODNP + YODNP*YODNP + ZODNP*ZODNP)
       DIFLA = DODNP - RGC
       IF (ABS (DIFLA) - 1.0E-5) 120, 120, 110
 110 ZDEDNP = ZDEDNP - DIFLA
 4001 FORMAT (1H , 5X, 'ODC 0, 0, 'F7.5, ' = GC X, Y, Z OF'3F8.5,
           DODNP = 'F9.5,' DIF OF 'F9.6, ' AT LOND LAT'F10.4, F8.4)
       GO TO 100
  120 CONTINUE
       PHINOF = ATAN2 (YODNP, XODNP) *RAD
       IF (PHINOF.LT.0.0)
                         PHINOF = PHINOF + 360.0
       TNOF = -ACOS(ZODNP/DODNP)*RAD + 90.0
       IF (NOPT.EQ.1) PRINT 4001, ZDEDNP, XODNP, YODNP, ZODNP, DODNP, DIFLA,
     1 PHINOF, TNOF
       SGCLATD = SIN(GCLATD/RAD)
       CGCLATD = COS(GCLATD/RAD)
       SGCLOND = SIN(GCLOND/RAD)
       CGCLOND = COS(GCLOND/RAD)
C
          GET GEOCENTRIC X Y Z COORDINATES
       XGC = RGC*CGCLATD*CGCLOND
       YGC = RGC*CGCLATD*SGCLOND
       ZGC = RGC*SGCLATD
       GCT = (90.0 - GCLATD)/RAD
       SGCT = SIN(GCT)
       CGCT = COS(GCT)
С
          FIND X Y Z IN LOCAL COORDINATES OF X=0, Y=0, Z
С
                    THE LOCAL COORDINATE Z AXIS PASSES THRU P
C
                    THE LOCAL COORDINATE X, Z PLANE CONTAINS P
```

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GCROT = ATAN2 (YGC, XGC)
       IF (NOPT.EQ.1) PRINT 2001, XGC, YGC, ZGC, GCROT
 2001 FORMAT (1H , 4F10.4,
                                3X, 'XGC, YGC, ZGC, GCROT')
       SGCROT = SIN(GCROT)
       CGCROT = COS(GCROT)
       XRL = XGC*CGCROT*CGCT + YGC*SGCROT*CGCT - ZGC*SGCT
       YRL = -XGC*SGCROT + YGC*CGCROT
       ZRL = XGC*CGCROT*SGCT + YGC*SGCROT*SGCT + ZGC*CGCT
 2002 FORMAT (1H , 3F10.4,
                                 13X, 'XRL, YRL, ZRL')
       IF (NOPT.EQ.1) PRINT 2002, XRL, YRL, ZRL
        DETERMINE LOCATION OF OFFSET DIPOLE CENTER IN THESE SAME
                               ROTATED LOCAL COORDINATES
C
      XEDRL = XEDFGC*CGCROT*CGCT + YEDFGC*SGCROT*CGCT - ZEDFGC*SGCT
       YEDRL = -XEDFGC*SGCROT + YEDFGC*CGCROT
       ZEDRL = XEDFGC*CGCROT*SGCT + YEDFGC*SGCROT*SGCT + ZEDFGC*CGCT
       IF (NOPT.EQ.1) PRINT 3003, XEDRL, YEDRL, ZEDRL
 3003 FORMAT (1H , 3F10.4, 13X, 'XEDRLM YEDRL, ZEDRL')
      XEDRTL = XEDRL
       YEDRTL = YEDRL
       ZEDRTL = ZEDRL - ZRL
       IF (NOPT.EQ.1) PRINT 2303, XEDRTL, YEDRTL, ZEDRTL
 2303 FORMAT (1H , 3F10.4, 13X, 'XEDRTL, YEDRTK, ZEDRTL')
          TRANSLATE TO LOCAL COORDINATE SYSTEM WITH ORIGIN AT SURFACE
      XRTL = XRL
      YRTL = YRL
      ZRTL = -ZRL
      XEDP = XRTL + XEDRL
      YEDP = YRTL + YEDRL
      ZEDP = ZRTL + ZEDRL
      RED = SQRT (XEDP*XEDP+YEDP*YEDP+ZEDP*ZEDP)
2302 FORMAT (1H , 3F10.4, 13X, 'XRTL, YRTL, ZRTL')
      IF (NOPT.EQ.1) PRINT 2302, XRTL, YRTL, ZRTL
                                 EARTHS SURFACE AT A SPECIFIED ALTITUDE
      POSITION OF OFFSET NORTH DIPOLE IN LOCAL COORDINATE SYSTEM
      XODNPR= XODNP*CGCROT*CGCT + YODNP*SGCROT*CGCT - ZODNP*SGCT
      YODNPR= -XODNP*SGCROT + YODNP*CGCROT
               XODNP*CGCROT*SGCT + YODNP*SGCROT*SGCT + ZODNP*CGCT
      ZODNPR=
      XODNPT = XODNPR
      YODNPT = YODNPR
       ZODNPT = ZODNPR - ZRL
       IF (NOPT.EQ.1) PRINT 1103, XODNPT, YODNPT, ZODNPT
 1103 FORMAT(1H ,10X,'XODNPT ='F10.4,2X,'YODNPT ='F10.4,2X,'ZODNPT ='
     1 F10.4,3X, 'OFFSET N DIPOLE IN LOCAL COORDINATES')
       ROTM = ATAN2 (YODNPT, XODNPT) + PI
C
                    FIND ANGLE FROM GEOGRAPHIC NORTH
С
                    NEGATIVE - ROTATION FROM GEOGRAPHIC NP CLOCKWISE
C
                    POSITIVE - ROTATION FROM GEOGRAPHIC NP CCW
       SROTM = SIN(ROTM)
       CROTM = COS(ROTM)
       ROTMD = ROTM*RAD
 2327 FORMAT (1H , F15.5, 3X, ' ROTM IN DEGREES MEASURED CCW SO -X
     1 WILL POINT TOWARD OFFSET NORTH DIPOLE AXIS')
       IF (NOPT.EQ.1) PRINT 2327, ROTMD
C
          FIND COMPONENTS OF UNIT VECTOR AT ARBITARY AZIMUTH AND ZENITH
       PLAZ = -AZ/RAD + PI
       TLZE = ZE/RAD
       SPLAZ = SIN(PLAZ)
       CPLAZ = COS(PLAZ)
       STLZE = SIN(TLZE)
       CTLZE = COS(TLZE)
```

```
IF (NOPT.EQ.1) PRINT 2005, XLD, YLD, ZLD, AZ, ZE
    2005 FORMAT (1H , 5F10.4, 3X, 'UNIT VECTOR COMPOENTS AT AZ & ZE')
   C
            FIND COMPONENTS OF UNIT VECTOR IN DIPOLE RADIAL COORDINATES
   C
   C
           ROTATE AROUND Y AXIS SO -Z AXIS PASSES THROUGH XED, 0, ZED
   C
           NEW VECTOR IS VA = ZRTL + ZEDRTK + XEDRTL
   C
           ANGLE BETWEEN VECTOR FROM POINT LOCAL ORIGIN TO GEOCENTER
   C
           AND VECTOR FROM POINT LOCAL ORIGIN TO XED, 0, ZED
   JIM LANGWORTHY'S FIX
   CA=DPEC (XEDRTL, ZEDRTL, ZRTL)
           CA = ZRTL*ZEDRTL/(ABS(ZRTL)*SQRT(ZEDRTL*ZEDRTL + XEDRTL*XEDRTL))
          A = ACOS(CA)
          IF(XEDRTL.GT.0.0) A = -A
          SA = SIN(A)
          ADEG = A*RAD
          IF (NOPT.EQ.1) PRINT 1000, CA, A, SA, ADEG
          XLP = XLD*CA + ZLD*SA
                       YLD
          YLP =
          ZLP = -XLD*SA +
                            ZLD*CA
          IF (NOPT.EQ.1) PRINT 5001, XLP, YLP, ZLP
    5001 FORMAT(1H , 3F10.4, 13X, 'XLP, YLP, ZLP ')
           ROTATE AROUND X PRIME AXIS SO -Z PASSES THROUGH XED, YED, ZED
   C
           CB=DPEC (YEDRTL, ZEDRTL, ZRTL)
          CB = ZRTL*ZEDRTL/(ABS(ZRTL)*SQRT(ZEDRTL*ZEDRTL + YEDRTL*YEDRTL))
          B = ACOS(CB)
          IF(YEDRTL.GT.0.0) B = -B
          SB = SIN(B)
          BDEG = B*RAD
Ŀŵ
          IF (NOPT.EQ.1) PRINT 1000, CB, B, SB, BDEG
XLPP = XLP
          YLPP = YLP*CB + ZLP*SB
          ZLPP =-YLP*SB + ZLP*CB
          IF (NOPT.EQ.1) PRINT 5002, XLPP, YLPP, ZLPP
    5002 FORMAT(1H , 3F10.4, 13X, 'XLPP, YLPP, ZLPP ')
           ROTATE AROUND ZPP AXIS SO -X AXIS PASSES THROUGH NORTH
   C
                                            OFFSET DIPOLE AXIS
   C
          ZLDM = ZLPP
          XLDM = XLPP*CROTM + YLPP*SROTM
          XLDM = XLPP*CROTM -YLPP*SROTM
          YLDM =-XLPP*SROTM + YLPP*CROTM
          YLDM = XLPP*SROTM + YLPP*CROTM
          IF (NOPT.EQ.1) PRINT 1101, XLD, YLD, ZLD, XLDM, YLDM, ZLDM
    1101 FORMAT (1H , 'UNIT VECTOR IN LOCAL COORDINATES ', 3F8.5,5X,
        1 'UNIT VECTOR IN LOCAL MAGNETIC COORDINATES', 3F10.5)
             FIND AZUMITH ANGLE OF UNIT VECTOR IN LOCAL DIPOLAR RADIAL COOR
    C
          IF((ABS(YLDM).GT.SMALL).OR.(ABS(XLDM).GT.SMALL)) GO TO 1102
          PAZM=0.0
          GO TO 1104
    1102 PAZM = ATAN2 (YLDM, XLDM)
    1104 AZM = (PI - PAZM) *RAD
          IF(AZM.GT.360.0) AZM = AZM - 360.0
          ZEM = ACOS(ZLDM)*RAD
```

FIND GAMMA ANGLE

GAMMA = ACOS (YLDM) *RAD

XLD = STLZE*CPLAZ
YLD = STLZE*SPLAZ

ZLD = CTLZE

```
C
       XED1=XGC-XEDFGC
       YED1=YGC-YEDFGC
       ZED1=ZGC-ZEDFGC
       FIND THE Z COORDINATE IN OFFSET DIPOLE COORDINATES
C
       ZED2=XED1*CPST-YED1*SPST+ZED1*CT
       FIND THE GEOMAGNETIC LATITUDE
C
       EDLAT=RAD* (PIO2-ACOS (ZED2/RED))
       COSLDA=COS (EDLAT/RAD)
       STORMER=60.*COSLDA**4./
               (RED*RED* (1.+SQRT (1.-COSLDA**3.*YLDM)) **2)
      RETURN
      END
C-----
     REAL FUNCTION DPEC (U, ZEDRTL, ZRTL)
     IMPLICIT NONE
     REAL U, ZEDRTL, ZRTL
                  ______
     DPEC=SIGN(1./SNGL(DSQRT(1D0+DBLE((U/ZEDRTL)**2);), ZRTL*ZEDRTL)
     RETURN
     END
      SUBROUTINE GridApproxLval(Cgrid, XLbounds, ILbins, Grid80Lval,
                    UseLapprox)
      IMPLICIT NONE
      REAL XLinfinite, RatioCheck, Grid80Lval, Cgrid
      INTEGER ILmax, NLvals, L, ILbins
      PARAMETER (XLinfinite=1.0E+06, NLvals=10, RatioCheck=1.2)
      REAL XLbounds (NLvals)
      LOGICAL UseLapprox
       Grid80Lval=XLinfinite
       IF (Cgrid .GT. 0.) Grid80Lval=SQRT(14.5/Cgrid)
       ILmax = ILbins
       UseLapprox=.FALSE.
       IF (Grid80Lval .GT. RatioCheck*XLbounds(ILmax)) THEN
         UseLapprox=.TRUE.
       ELSE
         DO L=2, ILmax
          IF ( (Grid80Lval .GT. RatioCheck*XLbounds(L-1)) .AND.
             (Grid80Lval .LT. XLbounds(L)/RatioCheck) ) THEN
              UseLapprox=.TRUE.
```

TRANSFORM TO OFFSET DIPOLE COORDINATES

ENDIF ENDDO ENDIF

RETURN END

END

```
SUBROUTINE GET_BLCOORDS (YEAR, LATI, LONGI, HEIGHT, XL, BB0, ICODE)
C
        Subroutine -- adapted from BILCAL by AJT -- for calculating
C
        geomagnetic coordinates B/B0 (=BB0) and McIlwain L (=XL)
С
        12/9/92
C
С
        Modified 11-17-97: add IMPLICIT NONE & variable-type declarations
C
С
        Inputs:
C
        YEAR = year (eg., 1987.63) for field initialization, etc.
C
        LATI, LONGI = geodetic latitude and (east) longitude (degrees)
С
                    = geodetic altitude (km above sea level)
        HEIGHT
C
        Outputs:
C
        XL = McIlwain L parameter
C
        BB0 = B/B0
C
        ICODE = return code: 1=OK; 3=approx result;
C
                              2=one of the conjugate mirror points is
C
                                 unphysical.
        IMPLICIT NONE
        REAL YEAR, LATI, LONGI, HEIGHT, XL, BBO
        INTEGER ICODE
        LOGICAL VAL
        REAL DIMO, BNORTH, BEAST, BDOWN, BABS, BAB1, BEQU, BDEL, BEQ, RRO
        Initialize field coefficients (if needed), get dipole moment
        CALL FELDCOF (YEAR, DIMO)
        Get local field strength (BABS)
        CALL FELDG (LATI, LONGI, HEIGHT, BNORTH, BEAST, BDOWN, BABS)
        Calculate McIlwain L and set ICODE flag.
        CALL SHELLG (LATI, LONGI, HEIGHT, DIMO, XL, ICODE, BAB1)
        IF (IABS (ICODE) .GT.9) ICODE=2
С
        Calculate B/B0
        BEQU=DIMO/(XL*XL*XL)
        IF (ICODE.EQ.1) THEN
                BDEL=1.E-3
                 CALL FINDBO (0.05, BDEL, VAL, BEQ, RRO)
                 IF(VAL) BEQU=BEQ
        ENDIF
        BB0=BABS/BEQU
        IF(BB0.GT.9999.999) BB0=9999.999
C
        Done
        RETURN
```

SUBROUTINE GET_CHECK_CONTROL(FILE_CHECK)

Sets logical to direct various file-checking functions in the CREME96 software (CHECK_FILE, CHECK_NAME_CONFLICT, CHECK_OUTPUT_FILE)

Recommended usage: logical FILE_CHECK=.true.

for VAX & PC stand-alone versions of the code

logical FILE_CHECK=.false.
for the PC/WWW version.

IMPLICIT NONE LOGICAL FILE_CHECK

FILE_CHECK=.true.

RETURN END

C C

С С С

C

C C

С

```
REAL FUNCTION GET_CREME96_FLUX(IZ, EN, YEAR, IMODE, ITRANS)
С
С
        Returns particle flux from CREME96 particle environment model.
        IMPLICIT NONE
        INTEGER*4 IZ, IMODE, ITRANS
        REAL*4 EN, YEAR, FLUX
        REAL*4 CRF96, CUT96, GET_TRAPPED_PROTONS, GET_TRAPPED_IONS
        GET_CREME96_FLUX=0.0
        IF (EN.LT.1.0) RETURN
        IF (IZ.LT.1 .or. IZ.GT.92) RETURN
        IF (ITRANS.EQ.0) THEN
C
            Fluxes outside of the magnetosphere
            FLUX=CRF96 (IZ, EN, YEAR, IMODE)
        ELSEIF (ITRANS.EQ.1) THEN
C
            Non-trapped fluxes inside the magnetosphre
            FLUX=CUT96 (IZ, EN, YEAR, IMODE)
        ELSEIF (ITRANS.EQ.2) THEN
C
            Non-trapped & Trapped fluxes inside the magnetosphere
            FLUX=CUT96 (IZ, EN, YEAR, IMODE)
C
            Function names modified 12-10-97 by AJT
            IF (IZ.EQ.1) FLUX=FLUX+GET_TRAPPED_PROTONS(EN)
            IF (IZ.GT.1) FLUX=FLUX+GET_TRAPPED IONS(IZ,EN)
        ENDIF
        GET_CREME96_FLUX=FLUX
        RETURN
        END
```

```
C
          Decodes header of CREME96 flux file to determine type of flux model.
   С
          Information is used in converting solar particle results from average
   С
          rates to event-accumulated numbers.
   C
   C
          IMPLICIT NONE
          CHARACTER*80 INFILE, ILINE
          CHARACTER*3 SUFFIX, IMODEL
           INTEGER*4 MODEL TYPE, ILONG, ILONG1, ILONG2,
                     IVER, J, NHEADER, STAT, CREME 96 OPEN
          MODEL_TYPE=0
           ILONG=INDEX(inFILE,'.')
           SUFFIX=INFILE(ILONG+1:ILONG+3)
          CALL CAPITALIZE_STRING(SUFFIX, 3)
           IF (SUFFIX.ne.'FLX' .and. SUFFIX.ne.'TFX' .and.
               SUFFIX.ne.'LET' .and. SUFFIX.ne.'DLT') RETURN
   С
   С
          Now open file and decode header:
   С
           CALL CHECK_CREME96_VERSION(INFILE, IVER)
           IF (IVER.GE.102) THEN
               stat = creme96_open(infile,'user',10,'old')
               IF (STAT.EQ.O) THEN
               READ(10,*) NHEADER
DO J=1, NHEADER
                  READ(10,110) ILINE
14
                  FORMAT (A80)
    110
3
                  IMODEL='
                  ILONG1=INDEX(ILINE,'%IMODE =')
1 74
                  IF (ILONG1.NE.0) IMODEL=ILINE(ILONG1+8:ILONG1+10)
IJ
                  ILONG2=INDEX(ILINE,'%IMODE =')
                  IF (ILONG2.NE.0) IMODEL=ILINE(ILONG2+10:ILONG2+12)
                                    ') THEN
                  IF (IMODEL.NE.'
                       WRITE(6,9999) IMODEL
                       FORMAT (' IMODEL=', A3, '====')
    C 9999
                      DECODE (3,100, IMODEL) MODEL TYPE
                      FORMAT(I3)
    100
                      RETURN
                  ENDIF
                ENDDO
                ENDIF
           ENDIF
           RETURN
           END
```

SUBROUTINE GET_CREME96_FLUX_MODEL(INFILE, MODEL_TYPE)

	С	SUBROUTINE GET_CREME96_VERSION(IVER)
	C C	Sets version number of CREME96 software, for record keeping purposes:
		IMPLICIT NONE INTEGER*4 IVER
	С	Modified 7/29/96: Version 1.01
	С	Modified 8/19/96 Version 1.02; more extensive output file headers
	С С С	Modified 9/14/96 Version 1.03: default energy limits; energy limits in LET calculations extended tables for Z > 28 added.
	C	Modified 9/25/96 Version 1.04: ACR charge-state distributions added IVER=104
		RETURN END

```
REAL FUNCTION GET_GTF(RIGIDITY)
C
C
         Evaluates orbit-averaged geomagnetic transmission function
С
          (previously calculated by GEOMAG96 and loaded into COMMON/GTFDAT
C
         by LOAD_GTF) at rigidity RIGIDITY (in GV).
C
         IMPLICIT NONE
         INTEGER*4 NGTF, IGTF, I, ISAV
         REAL*4 R,GTF,P,RIGIDITY
         PARAMETER (NGTF=1001)
         COMMON/GTFDAT/IGTF, R (NGTF), GTF (NGTF)
C
С
         LOOK UP THE TABULATED MAGNETIC RIGIDITY JUST ABOVE RIGID
C
         GET GTF=0.0
         IF (IGTF.LE.0) RETURN
         P=RIGIDITY
         GET GTF=GTF(IGTF)
         IF (P.GT.R(IGTF)) RETURN
         GET GTF=0.0
         IF (P.LT.R(1)) RETURN
         DO 2 I=2, IGTF
         IF(P.GT.R(I)) GOTO 2
         ISAV=I
         GOTO 3
 2
         CONTINUE
С
С
         INTERPOLATE THE TRANSMISSION FACTOR (AVERAGED FOR THE ORBIT).
C
3
         GET GTF=GTF(ISAV-1)+
          (\mathtt{GTF}(\mathtt{ISAV}) - \mathtt{GTF}(\mathtt{ISAV-1})) \star (\mathtt{P-R}(\mathtt{ISAV-1})) / (\mathtt{R}(\mathtt{ISAV}) - \mathtt{R}(\mathtt{ISAV-1}))
         RETURN
         END
```

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```
SUBROUTINE GET_HI_XS_INPUTS (DEVICE LABEL, XM, YM, ZM, FUNNELM, NBITS,
     &
                                     IPARAM, PARAMS, XSECT FILE)
C
C
        Interactive dialogue to get necessary cross-sectioninput parameters
С
С
        for heavy-ion upsets:
C
C
        Written by:
                       Allan J. Tylka
C
                       Code 7654
C
                       Naval Research Laboratory
C
                       Washington, DC 20375-5352
C
                       tylka@crs2.nrl.navy.mil
C
C
        IMPLICIT NONE
        CHARACTER*80 XSECT FILE
        CHARACTER*40 DEVICE LABEL
        REAL*4 XM, YM, ZM, FUNNELM, PARAMS, NBITS
        INTEGER*4 IPARAM
        DIMENSION PARAMS (4)
        INTEGER*4 IFILETYPE, IACCEPT
        INTEGER*4 IERR
        DATA IERR/0/
       WRITE(6,1210)
1210 FORMAT(1x,' *** NOTE ***: At any point in the following'
                  ' dialogue, you can go back and'
    &
    &
             /,1x,' change a parameter by entering -1 for the',
                  ' presently requested information.',
    &
             /,lx,' Repeated -1 values can be used to scroll back',
     δ.
                  ' to (almost) anywhere in the input menu.')
105 CONTINUE
        CALL RETRY INPUT (IERR)
        WRITE (6, 1215)
1215
       FORMAT(/,1x,' Enter device label and/or comments',
               ' (40 characters max) for record-keeping:')
       READ(*,1218,ERR=105,IOSTAT=IERR) DEVICE LABEL
1218
       FORMAT (A40)
       IF (DEVICE LABEL (1:2).EO.'-1') RETURN
       WRITE(6,1219) DEVICE LABEL
1219
       FORMAT(1x,' Device/Comment: ',A40)
1185
       CONTINUE
       CALL RETRY INPUT (IERR)
       WRITE(6,1200)
1200 FORMAT(/,' This program calculates the heavy-ion SEU rate',
    &
                 ' using the RPP method.',
                /,' Enter the dimensions of the',
                  ' bit sensitive volume: (X,Y,Z; in microns)')
       READ(*, *, ERR=1185, IOSTAT=IERR) XM, YM, ZM
       IF (XM.LE.-1.0 .or. YM.LE.-1.0 .or. ZM.LE.-1.0) GOTO 105
       WRITE(6,1220) XM, YM, ZM
      FORMAT(' Sensitive volume dimensions = ',
1220
                F8.2, ' x ', F8.2, ' x ', F8.2, ' microns')
```

```
CONTINUE
1195
       CALL RETRY INPUT (IERR)
       WRITE (6, 2200)
       FORMAT (' Enter funnel length (microns): ')
2200
       READ (*, *, ERR=1195, IOSTAT=IERR) FUNNELM
       IF (FUNNELM.LE.-1.0) GOTO 1185
       WRITE(6,2220) FUNNELM
       FORMAT(' Funnel length = ',F8.2,' microns.')
2220
1225
       CONTINUE
       CALL RETRY INPUT (IERR)
       WRITE (6, 1300)
       FORMAT(//,' This code supports several methods for specifying',
1300
                  ' the SEU cross-section:',
               /,' METHOD = 0: a file containing a two-column table,'
               /,' METHOD = 1: Bendel 1-parameter fit ',
               /,' METHOD = 2: Bendel 2-parameter fit ',
               /,' METHOD = 3: NOT CURRENTLY USED',
              /,' METHOD = 4: Weibull fit',
               /, ' METHOD = 5: Critical charge (in pC)',
              //,' Specify METHOD (0,1,2,4, or 5): ')
       READ(*,*,ERR=1225,IOSTAT=IERR) IPARAM
       IF (IPARAM.LE.-1) GOTO 1195
       IF (IPARAM.EQ.0) THEN
1395
          CONTINUE
           CALL RETRY INPUT (IERR)
          WRITE (6, 1400)
          FORMAT(' SEU cross-section from input table file:',
               /,' This table must have a two-column format with :',
      /,11x,
               ' column 1 containing LET (in MeV-cm2/milligram)',
    &
      /,7x,' and column 2 containing SEU cross-section',
             ' (in sq. microns/bit)',
    & /,7x,' and be ordered according to increasing LET.',
      /,' The file containing the table must already exist in',
    &
      ' your current USER area and ',
    & /,' be called something.XSD (ie., have XSD for the extension).',
      /,' Enter name of the cross-section file: ')
       READ(*,1,ERR=1395,IOSTAT=IERR) XSECT FILE
1
       FORMAT (A80)
       IF (XSECT FILE(1:2).EQ.'-1') GOTO 1225
       WRITE (6,1410) XSECT FILE
1410
       FORMAT(1x,' Input Heavy-Ion Cross-Section File = ',/,1x,A80)
       IFILETYPE=8
       CALL CHECK FILE (IFILETYPE, XSECT_FILE, IACCEPT)
       IF (IACCEPT.NE.0) GOTO 1395
       ELSEIF (IPARAM.EQ.1) THEN
1495
           CONTINUE
           CALL RETRY_INPUT(IERR)
          WRITE(6,1500)
          FORMAT(' Bendel 1-parameter fit to the cross-section: ',
1500
                /,' NOTE: Your fit parameters must specify heavy-ion',
    æ
                /,' SEU cross-section (in sq. microns/bit) vs.',
                ' LET (in MeV-cm2/milligram): '
                /,' Enter Bendel-1 parameter value: ')
           READ(*,*,ERR=1495,IOSTAT=IERR) PARAMS(1)
```

```
IF (PARAMS(1).LE.-1.) GOTO 1225
           WRITE(6,1510) PARAMS(1)
1510
           FORMAT(' Bendel-1 parameter = ',E13.6)
       ELSEIF (IPARAM.EQ.2) THEN
           CONTINUE
1595
           CALL RETRY INPUT (IERR)
           WRITE (6, 1600)
           FORMAT(' Bendel 2-parameter fit to the cross-section: ',
1600
                    NOTE: Your fit parameters must specify heavy-ion',
                 /,' SEU cross-section (in sq. microns/bit) vs.',
    &
                  ' LET (in MeV-cm2/milligram): '
    &
                /,' Enter Bendel A & B parameter values: ')
    &
           READ(*,*,ERR=1595,IOSTAT=IERR) PARAMS(1),PARAMS(2)
           IF (PARAMS(1).LE.-1. .or. PARAMS(2).LE.-1.) GOTO 1225
           WRITE(6,1610) PARAMS(1), PARAMS(2)
           FORMAT(' Bendel parameters A, B = ',2E13.6)
1610
       ELSEIF (IPARAM.EQ.3.or.IPARAM.LT.0 .or. IPARAM.GT.5) THEN
           CONTINUE
1695
           WRITE(6,1700)
1700
           FORMAT (' ILLEGAL CROSS-SECTION SPECIFICATION CODE. ',
                /,' Please try again.')
           GOTO 1225
       ELSEIF (IPARAM.EO.4) THEN
1795
           CONTINUE
           CALL RETRY INPUT (IERR)
           WRITE(6,1800)
1800
           FORMAT(' Weibull fit to the cross-section: ',
                /,' NOTE: Your fit parameters must specify heavy-ion'
                /,' SEU cross-section (in sq. microns/bit) vs.',
    &
                 ' LET (in MeV-cm2/milligram): ')
    &
           WRITE(6,1810)
1810
           FORMAT(' Enter ONSET parameter (in MeV-cm2/milligram): ')
           READ(*,*,ERR=1795,IOSTAT=IERR) PARAMS(1)
           IF (PARAMS(1).LE.-1.) GOTO 1225
1815
           CONTINUE
           CALL RETRY INPUT (IERR)
           WRITE(6,1820)
           FORMAT(' Enter WIDTH parameter (in MeV-cm2/milligram): ')
1820
           READ (*, *, ERR=1815, IOSTAT=IERR) PARAMS (2)
           IF (PARAMS(2).LE.-1.) GOTO 1795
1825
           CONTINUE
           CALL RETRY INPUT (IERR)
           WRITE(6,1830)
1830
           FORMAT(' Enter POWER parameter (dimensionless exponent): ')
           READ(*,*,ERR=1825,IOSTAT=IERR) PARAMS(3)
           IF (PARAMS(3).LE.-1.) GOTO 1815
1835
           CONTINUE
           CALL RETRY INPUT (IERR)
           WRITE(6,1840)
           IF (XM.GT.0.0 .and. YM.GT.0.0) WRITE(6,1841)
1840
           FORMAT(' Enter cross-section plateau value',
                  ' (in sq. microns/bit).')
    &
           FORMAT(' (If 0, calculation will use surface area (xy) of',
1841
                  ' the RPP sensitive volume.)')
           READ(*,*,ERR=1835,IOSTAT=IERR) PARAMS(4)
           IF (PARAMS (4).LE.-1.) GOTO 1825
           IF (PARAMS(4).LE.O.) PARAMS(4) = XM*YM
```

```
WRITE(6,1850) PARAMS(1), PARAMS(2), PARAMS(3), PARAMS(4)
           FORMAT(' Weibull fit parameters: ',
1850
                 /,5x, ONSET = ',F9.3,' MeV-cm2/milligram',
    &
                 /,5x,' WIDTH = ',F9.3,' MeV-cm2/milligram',
                 /,5x,' POWER = ',F9.3,' (dimensionless)',
    &
                 /,5x,' PLATEAU = ',F9.3,' square microns/bit')
       ELSEIF (IPARAM.EQ.5) THEN
1895
           CONTINUE
           WRITE(6,1900)
1900
           FORMAT(' Cross-section given as step function in critical',
    &
                  ' charge.',
           /,' NOTE: in general this method does NOT'
    &
            ' give accurate results for space',
    &
           /,' applications, but it may be useful for'
             ' order-of-magnitude estimates by chip',/,' designers.')
1905
           CONTINUE
           CALL RETRY INPUT (IERR)
           WRITE(6,1910)
           FORMAT(' Enter critical charge (in picocoloubs): ')
1910
           READ(*, *, ERR=1905, IOSTAT=IERR) PARAMS(1)
           IF (PARAMS(1).LE.-1.) GOTO 1225
1915
           CONTINUE
           CALL RETRY INPUT (IERR)
           WRITE(6,1920)
           IF (XM.GT.0.0 .and. YM.GT.0.0) WRITE(6,1841)
1920
           FORMAT(' Enter cross-section (in square microns/bit):')
           READ(*,*,ERR=1915,IOSTAT=IERR) PARAMS(2)
           IF (PARAMS(2).LE.-1.) GOTO 1905
           IF (PARAMS(2).LE.O.) PARAMS(2) = XM*YM
           WRITE(6,1930) PARAMS(1), PARAMS(2)
           FORMAT(' Critical charge = ',E13.5,' picocoloumbs',
1930
                /,' Cross-Section = ',E13.5,' square microns/bit')
    &
        ENDIF
1995
        CONTINUE
        CALL RETRY INPUT (IERR)
        WRITE(6,2000)
2000
        FORMAT(' Finally, specify number of bits per device: ')
        READ (*, *, ERR=1995, IOSTAT=IERR) NBITS
        IF (NBITS.EQ.-1) THEN
           IF (IPARAM.EQ.1) GOTO 1495
           IF (IPARAM.EQ.2) GOTO 1595
           IF (IPARAM.EQ.3) GOTO 1695
           IF (IPARAM.EQ.4) GOTO 1835
           IF (IPARAM.EQ.5) GOTO 1915
        ENDIF
        WRITE(6,2010) NBITS
2010
        FORMAT(1x,E13.5,' bits per device.')
        RETURN
        END
```

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```
SUBROUTINE GET PROTON XS INPUTS (DEVICE LABEL, NBITS,
     &
                                        IPARAM, PARAMS, XSECT FILE)
C
C
C
        Generates interactive dialogue to get necessary input parameters
С
        for proton-induced SEU rate:
С
C
       Written by:
                       Allan J. Tylka
С
                       Code 7654
C
                       Naval Research Laboratory
C
                       Washington, DC 20375-5352
С
                       tylka@crs2.nrl.navy.mil
C
C
       Last update:
                     20 August 1996
C
C-
С
       IMPLICIT NONE
       CHARACTER*80 XSECT FILE
       CHARACTER*40 DEVICE LABEL
       REAL*4 PARAMS, NBITS
       INTEGER*4 IPARAM
       DIMENSION PARAMS (4)
       INTEGER*4 IFILETYPE, IACCEPT, IERR
       DATA IERR/0/
       WRITE(6,1210)
1210 FORMAT(1x,' *** NOTE ***: At any point in the following'
                  ' dialogue, you can go back and'
    &
           /,1x,' change a parameter by entering -1 for the',
                 ' presently requested information.',
    ₽.
            /,lx,' Repeated -1 values can be used to scroll back',
                  ' to (almost) anywhere in the input menu.')
 105
       CONTINUE
       CALL RETRY_INPUT(IERR)
       WRITE(6,1215)
1215
       FORMAT(/,1x,' Enter device label and/or comments',
              ' (40 characters max) for record-keeping:')
       READ(*,1218,ERR=105,IOSTAT=IERR) DEVICE LABEL
1218
       FORMAT (A40)
       IF (DEVICE LABEL(1:2).EQ.'-1') RETURN
       WRITE(6,1219) DEVICE LABEL
1219
       FORMAT(1x,' Device/Comment: ',A40)
1295
       CONTINUE
       CALL RETRY INPUT (IERR)
       WRITE(6,1300)
       FORMAT( /,' This code supports several methods for specifying',
1300
                  ' the SEU cross-section:',
               /,' METHOD = 0: a file containing a two-column table,'
               /,' METHOD = 1: Bendel 1-parameter fit',
               /,' METHOD = 2: Bendel 2-parameter fit ',
               /,' METHOD = 3: NOT CURRENTLY USED',
               /,' METHOD = 4: Weibull fit',
              //,' Specify METHOD (0,1,2, or 4): ')
```

```
READ(*,*,ERR=1295,IOSTAT=IERR) IPARAM
        IF (IPARAM.EQ.-1) GOTO 105
        IF (IPARAM.EQ.0) THEN
 1395
            CONTINUE
            CALL RETRY INPUT (IERR)
            WRITE(6,1400)
            FORMAT(' SEU cross-section from input table file:',
 1400
                 /,' This table must have a two-column format with :',
     &
                 ' column 1 containing proton energy (in MeV)',
     &
        /,7x,' and column 2 containing SEU cross-section',
              ' (in 10**-12 cm2/bit)',
       /,7x,^{\prime} and be ordered according to increasing proton energy.^{\prime},
        /,' The file containing the table must already exist in',
           ' your current USER area and ',
         /,' be called something.XSD (ie., have XSD for the extension).',
        /,' Enter name of the cross-section file: ')
        READ(*,1,ERR=1395,IOSTAT=IERR) XSECT_FILE
1
        FORMAT (A80)
        IF (XSECT_FILE(1:2).EQ.'-1') GOTO 1295
        WRITE(6,1410) XSECT FILE
1410
       FORMAT(1x,' Input Proton Cross-Section File = ',/,1x,A80)
        IFILETYPE=8
       CALL CHECK FILE (IFILETYPE, XSECT_FILE, IACCEPT)
       IF (IACCEPT.NE.0) GOTO 1395
       ELSEIF (IPARAM.EQ.1) THEN
1495
           CONTINUE
            CALL RETRY INPUT (IERR)
            WRITE (6, 1500)
           FORMAT(' Bendel 1-parameter fit to the cross-section: ',
1500
                 /,' NOTE: Your fit parameters must specify proton',
    &
                 /,' SEU cross-section (in 10**-12 cm2/bit) vs.',
                  ' proton energy (in MeV): '
    &
                 /,' Enter Bendel-1 parameter value: ')
           READ(*, *, ERR=1495, IOSTAT=IERR) PARAMS(1)
           IF (PARAMS(1).LE.-1) GOTO 1295
           WRITE(6,1510) PARAMS(1)
1510
           FORMAT(' Bendel-1 parameter = ',E13.6)
       ELSEIF (IPARAM.EQ.2) THEN
1595
           CONTINUE
           CALL RETRY INPUT (IERR)
           WRITE(6,1600)
           FORMAT(' Bendel 2-parameter fit to the cross-section: ',
1600
           /,' NOTE: Your fit parameters must specify the proton',
    æ
           /,' SEU cross-section (in 10**~12 cm2/bit) vs.',
            ' proton energy (in MeV): '
           /,' Enter Bendel A & B parameter values: ')
           READ(*,*,ERR=1595,IOSTAT=IERR) PARAMS(1),PARAMS(2)
           IF (PARAMS(1).LE.-1. .or. PARAMS(2).LE.-1.) GOTO 1295
           WRITE(6,1610) PARAMS(1), PARAMS(2)
           FORMAT(' Bendel parameters A,B = ',2E13.6)
1610
       ELSEIF (IPARAM.EQ.3.or.IPARAM.LT.0 .or. IPARAM.GT.4) THEN
1695
           CONTINUE
           WRITE (6, 1700)
1700
           FORMAT(' ILLEGAL CROSS-SECTION SPECIFICATION CODE. ',
                /,' Please try again.')
```

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```
ELSEIF (IPARAM.EQ.4) THEN
1795
           CONTINUE
           CALL RETRY INPUT (IERR)
           WRITE (6, 1800)
           FORMAT(' Weibull fit to the cross-section: ',
1800
           1."
                 NOTE: Your fit parameters must specify the proton'
    &
                SEU cross-section (in 10**-12 cm2/bit) vs.',
            ' proton energy (in MeV): ')
           WRITE(6,1810)
           FORMAT(' Enter ONSET parameter (in MeV): ')
1810
           READ(*, *, ERR=1795, IOSTAT=IERR) PARAMS(1)
           IF (PARAMS(1).LE.-1.) GOTO 1295
1815
           CONTINUE
           CALL RETRY INPUT (IERR)
           WRITE(6,1820)
           FORMAT (' Enter WIDTH parameter (in MeV): ')
1820
           READ (*, *, ERR=1815, IOSTAT=IERR) PARAMS (2)
           IF (PARAMS(2).LE.-1.) GOTO 1795
1825
           CONTINUE
           CALL RETRY INPUT (IERR)
           WRITE (6, 1830)
1830
           FORMAT(' Enter POWER parameter (dimensionless exponent): ')
           READ (*, *, ERR=1825, IOSTAT=IERR) PARAMS (3)
           IF (PARAMS(3).LE.-1.) GOTO 1815
1835
           CONTINUE
           CALL RETRY_INPUT(IERR)
           WRITE(6,1840)
           FORMAT(' Enter cross-section plateau value',
1840
                  ' (in 10**-12 cm2/bit):')
           READ(*,*,ERR=1835,IOSTAT=IERR) PARAMS(4)
           IF (PARAMS (4).LE.-1.) GOTO 1825
           WRITE(6,1850) PARAMS(1), PARAMS(2), PARAMS(3), PARAMS(4)
1850
           FORMAT(' Weibull fit parameters: ',
                 /,5x,' ONSET = ',F9.3,' MeV',
    &
                                = ',F9.3,' MeV',
                 /,5x,' WIDTH
    &
                 /,5x,' POWER = ',F9.3,' (dimensionless)',
    ۶
                 /,5x, PLATEAU = ',F9.3,' x 10**-12 cm2/bit')
        ENDIF
1995
        CONTINUE
        CALL RETRY INPUT (IERR)
        WRITE (6, 2000)
2000
        FORMAT(' Finally, specify number of bits per device: ')
        READ(*, *, ERR=1995, IOSTAT=IERR) NBITS
        IF (NBITS.EQ.-1) THEN
           IF (IPARAM.EQ.1) GOTO 1495
           IF (IPARAM.EQ.2) GOTO 1595
           IF (IPARAM.EQ.3) GOTO 1695
           IF (IPARAM.EQ.4) GOTO 1835
        ENDIF
        WRITE(6,2010) NBITS
        FORMAT(1x,E13.5,' bits per device.')
2010
        RETURN
        END
```

С

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REAL FUNCTION GET_TRAPPED_IONS(IZ,EN)

Returns orbit-averaged flux of trapped ion IZ at energy EN --> NOT INCLUDED IN CREME96

GET_TRAPPED_IONS=0.0 RETURN END

```
REAL FUNCTION GET TRAPPED PROTONS (EN)
C
        Returns orbit-averaged trapped proton flux (in protons/m2-s-sr-MeV)
C
        at energy EN (in MeV) by interpolating value from previously stored
C
C
        array.
С
        Formerly called "TRAPPED PROTONS". Renamed by AJT 12-9-97, to
C
        remove name conflict with new PRB/SAIC routines on trapped protons.
С
C
C
        IMPLICIT NONE
        INTEGER*4 MAXSPEC, ITRPSPEC, I
        REAL*4 EN, ENTRP, FLUXTRP
        PARAMETER (MAXSPEC=5000)
        COMMON/TRPDAT/ITRPSPEC, ENTRP (MAXSPEC), FLUXTRP (MAXSPEC)
        REAL*4 X1, Y1, X2, Y2, X3, Y3, SLOPE
        GET_TRAPPED_PROTONS=0.
        IF (EN.LT.ENTRP(1) .or. EN.GT.ENTRP(ITRPSPEC)) RETURN
        DO 100 I=2, ITRPSPEC
                IF (EN.LE.ENTRP(I)) THEN
                    IF (FLUXTRP(I).GT.0. .and. FLUXTRP(I-1).GT.0.) THEN
                        X1=ALOG(ENTRP(I-1))
                        Y1=ALOG(FLUXTRP(I-1))
                        X2=ALOG(ENTRP(I))
                        Y2=ALOG(FLUXTRP(I))
                        SLOPE = (Y2 - Y1) / (X2 - X1)
                        X3=ALOG(EN)
                        Y3=SLOPE*(X3-X1)+Y1
                        GET TRAPPED PROTONS=EXP(Y3)
                        GOTO 150
                    ELSE
                        GET TRAPPED PROTONS=0.0
                    ENDIF
                ENDIF
 100
             CONTINUE
             CONTINUE
 150
        RETURN
        END
```

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C

SUBROUTINE GET_UPSET(XM,YM,ZM,FUNNELM,QCRIT, NVAL,LVAL,FVAL,UPSET)

Subroutine for evaluating SEUs as a function of critical charge. Derived from the UPSET program in the original CREME software.

Inputs: XM, YM, ZM = bit dimensions x, y, z (in microns)

FUNNELM = funnel length (in microns)

Qcrit = (critical charge) in picocoulombs

LVAL, FVAL = arrays containing integral LET spectrum

in particles/m2-sr-s vs. LET in MeV-cm2/gm

in NVAL points

Output: UPSET = SEUs/bit/s

Modified by AJT 10-24-96: increases density of sampling in LET spectrum around values corresponding to peaks in the differential pathlength distribution. This removes a previously noted problem in the old GET_UPSET routine, in which the SEU rate was not a strictly non-increasing function of increasing critical charge.

Modified by AJT 10-30-96: funnels added.

Modified by AJT 11-07-96: changed to trapezoidal rule integration, as suggested by Ed Petersen.

BANNER FROM CREME 1981 VERSION:

THIS PROGRAM COMPUTES THE UPSET RATE DUE TO THE
DIRECT IONIZATION OF INDIVIDUAL PARTICLES. IT ASSUMES
THAT FOR EACH BIT THERE EXISTS A SENSITIVE VOLUME.
IF AN AMOUNT OF ELECTRICAL CHARGE (>QCRIT) IS CREATED
WITHIN THIS VOLUME BY THE IONIZING PARTICLE, THEN AN
UPSET WILL OCCUR. THE SENSITIVE VOLUME IS IDEALIZED AS
A PARALLELEPIPED WITH DIMENSIONS X, Y, AND Z.
THE OUTPUT IS GIVEN IN UPSETS/(BIT*SECOND) AND UPSETS/(BIT*DAY).

THIS CALCULATION USES THE METHOD DESCRIBED IN "THE VARIABILITY OF SINGLE EVENT UPSET RATES IN THE NATURAL ENVIRONMENT,"

JAMES H. ADAMS, JR., IEEE TRANS. ON NUCL. SCI., NS-30, 4475-4480, DEC., 1983.

IT IS OPERATIONALLY EQUIVALENT TO ROCKWELL'S CRIER PROGRAM (PICKEL AND BLANDFORD, IEEE TRANS. ON NUCL. SCI. NS-26, DEC. 1979, PP.4735-4739) WHEN USED WITH HEINRICH'S LET SPECTRUM (W. HEINRICH, RADIATION EFFECTS, VOL. 34, PP. 143-8, 1977).

THIS PROGRAM CALLS A SUBPROGRAM, DIFPLD, THAT RETURNS
THE DIFFERENTAL PATHLENGTH DISTRIBUTION IN THE SENSTIVE VOLUME

IMPLICIT NONE

INTEGER*4 NBINS, NVAL, I, K, N

REAL*4 XM, YM, ZM, FUNNELM, QCRIT, LVAL, FVAL, UPSET

REAL*4 L, FLUX, X, Y, Z, DSI, AREA, PMAX, ENERGY, LMIN, SUM, Q

REAL*4 D, DIFPLD

REAL*4 FUNNEL

REAL*4 INTGRND0, INTGRND

PARAMETER (NBINS=5000)

```
DIMENSION L (NBINS), FLUX (NBINS)
        DIMENSION LVAL(1), FVAL(1)
        DATA DSI/2.321/
        REAL*4 LUSE, FUSE
        DIMENSION LUSE (NBINS), FUSE (NBINS)
        INTEGER*4 M, IENTER
С
        Load the inputs
        IF (IENTER.EQ.0) THEN
            IENTER=1
            WRITE (6, 9999)
 9999
            FORMAT(1X,' GET UPSET revision 11/08/96 active: ')
        ENDIF
        UPSET=0.0
C
        CALL TEMP STORAGE (NBINS, NVAL, LVAL, FVAL, N, L, FLUX)
C
С
        RPP Dimensions:
С
        X = XM
        Y=YM
        Z = ZM
        FUNNEL=FUNNELM
C
C
        CONVERT FROM MICROMETERS TO CENTIMETERS.
        X=X*.0001
        Y=Y*.0001
        Z=Z*.0001
        FUNNEL=FUNNEL*0.0001
С
C
        COMPUTE THE SURFACE AREA OF THE SENSITIVE VOLUME.
С
        AREA = (2.*X*Y+2.*X*Z+2.*Y*Z)
С
С
        CONVERT FROM SQUARE CENTIMETERS TO SQUARE METERS.
C
        AREA=AREA*.0001
C
С
        CONVERT THE DIMENSIONS OF THE SENSITIVE VOLUME TO G/CM**2.
C
        X=X*DSI
        Y=Y*DSI
        Z=Z*DSI
        FUNNEL=FUNNEL*DSI
С
С
        COMPUTE THE MAJOR DIAMETER OF THE SENSITIVE VOLUME.
С
        PMAX=SQRT(X*X+Y*Y+Z*Z)
C
C
        COMPUTE THE ENERGY (IN MEV) REQUIRED TO PRODUCE QCRIT(IN PC)
С
        HOLE-ELECTRON PAIRS IN SILICON.
С
        ENERGY=22.5*QCRIT
C
C
        COMPUTE THE MINIMUM LET THAT CAN PRODUCE AN UPSET.
C
С
        Funnel added to this equation 10/30/96 AJT
```

```
Now use expanded sampling in numerical integration:
    C
    C
             INTEGRATE FROM LMIN TO THE LARGEST LET IN THE SPECTRUM.
    C
     C
             SUM=0.0
             Q=LMIN
             Note: to uses the trapezoidal rule in the numerical integration,
     С
             we need to evaluate the integrand (DIFPLD*FLUX/L**2) on a grid
     C
             which includes the endpoints of the integration. The lower endpoint
     С
             is at LMIN, which corresponds to PMAX, the longest possible path
     C
             through the RPP. However, DIFPLD=0 at PMAX. Thus, the integrand
     С
             also vanishes at LMIN.
     С
             INTGRND0=0.0
The first that the first that the
             DO 10 I=1, M
             IF (LUSE(I).LT.LMIN) GO TO 10
     C
             Terminate numerical integration when integral flux falls to zero.
             IF (FUSE(I).LE.0.0) GOTO 11
132
     C
in in it
             COMPUTE THE PATHLENGTH CORRESPONDING TO L(I).
     C
     С
Funnel added to this equation 10/30/96 AJT
     C
ļ
             D=ENERGY/LUSE(I)
W.
             D= (ENERGY/LUSE(I) - FUNNEL)
             IF (D.LT.0.0) D=0.0
     С
             CARRY OUT THE INTEGRAL.
     C
             Modified to use trapezoidal rule 11/7/96:
             SUM=SUM+(LUSE(I)-Q)*DIFPLD(D,X,Y,Z)*FUSE(I)/(LUSE(I)**2)
     COLD
              INTGRND=DIFPLD(D,X,Y,Z)*FUSE(I)/(LUSE(I)**2)
              SUM=SUM+0.5*(LUSE(I)-Q)*(INTGRND+INTGRND0)
              INTGRND0=INTGRND
              Q=LUSE(I)
```

LMIN=ENERGY/PMAX

CONTINUE

CONTINUE

CONTINUE

COMPUTE THE ERROR RATE.

10

11 C C

12

C

LMIN=ENERGY/(PMAX+FUNNEL)

in the pathlength density distribution

Modification AJT 10/24/96: expand sampling around discontinuities

M, LUSE, FUSE)

CALL EXPAND SAMPLING (NBINS, N, L, FLUX, LMIN, ENERGY, PMAX, X, Y, Z,

С

С

C

C

C C &

```
RETURN
END
```

```
SUBROUTINE TEMP STORAGE (NBINS, NVAL, LVAL, FVAL, N, L, FLUX)
        IMPLICIT NONE
        INTEGER*4 NBINS, NVAL, N, K
        REAL*4 LVAL, FVAL, L, FLUX
        DIMENSION LVAL(1), FVAL(1), L(1), FLUX(1)
        Copy Integral LET spectrum:
С
        DO 5 K=1, NVAL
                  L(K) = LVAL(K)
                  FLUX(K)=FVAL(K)
        CONTINUE
 5
        N=NVAL
        IF (N.GT. NBINS) THEN
        WRITE(6,9999) N, NBINS
        FORMAT ('@ 10002 ABNORMAL TERMINATION: ',
 9999
          /,lx,' LET spec array out of bounds in GET_UPSET: ',
           /,1x, ' N = ', I8, ' NBINS = ', I8,
          /,1x,' STOP.')
        STOP
        ENDIF
        RETURN
        END
         SUBROUTINE EXPAND SAMPLING (NBINS, N, L, FLUX,
                                      LMIN, ENERGY, PMAX, X, Y, Z,
     &
                                      M, LUSE, FUSE)
     &
         IMPLICIT NONE
         INTEGER*4 NBINS, N, M, TBINS
         REAL*4 L, FLUX, LMIN, ENERGY, PMAX, X, Y, Z, LUSE, FUSE
         DIMENSION L(1), FLUX(1), LUSE(1), FUSE(1)
         PARAMETER (TBINS=5000)
         REAL*4 LTEMP, FTEMP
         DIMENSION LTEMP (TBINS), FTEMP (TBINS)
         INTEGER*4 I,K
         REAL*4 S, STEMP, SDUM, XVAL, YVAL
         REAL*4 SCALE, SMALLEST, SAMPLE SIZE
         DATA SCALE/0.01/
         REAL*4 DSI
        DATA DSI/2.321/
C
         For idiot checks:
         REAL*4 DUM, DIFPLD, STEMPDUM
C
         INTEGER*4 INDX
         DIMENSION INDX (TBINS)
```

```
NOTE: X,Y,Z assumed here to be in g/cm2!!!
C
C
       Store relevant portion of input LET spectrum:
C
       M = 0
       DO 4 I=1,N
          IF (L(I).LT.LMIN) GO TO 4
          IF (FLUX(I).LE.0.0) GOTO 6
          M=M+1
          LTEMP(M) = L(I)
          FTEMP(M)=FLUX(I)
       CONTINUE
 4
 6
       CONTINUE
       Now we wish to do additional samplings around the peaks
C
C
       in the pathlength distribution.
С
       Specifically, the additional sampling is done on a scale
       equal to 1% of the smallest dimension, but no larger than 0.01
C
C
       microns. The sampling is done at 100 points ranging from
       x-10*scale to x+90*scale.
C
       PMAX=SQRT(X*X+Y*Y+Z*Z)
       SMALLEST=MIN(X,Y,Z)/0.0001/DSI
       SAMPLE SIZE=SCALE*SMALLEST
        IF (SAMPLE SIZE.GT.0.01) SAMPLE SIZE=0.01
       DO 50 K=1,3
             IF (K.EQ.1) S=X
             IF (K.EQ.2) S=Y
             IF (K.EQ.3) S=Z
             Suppress redundant samplings:
C
             IF (K.EQ.2 .and. (ABS(X-Y).LE.0.0001*X)) GOTO 50
             IF (K.EQ.3 .and.
              (ABS(X-Z).LE.0.0001*X .or. ABS(Y-Z).LE.0.0001*Y)) GOTO 50
             DO 45 I=-11,89
                STEMP=S+FLOAT(I)*SAMPLE SIZE*0.0001*DSI
                IF (STEMP.LE.O. .or. STEMP.GT.PMAX) GOTO 45
C-----
С
                Idiot checks again:
C
                 DUM=DIFPLD(STEMP, X, Y, Z)
C
                 SDUM=S/DSI/0.0001
C
                 STEMPDUM=STEMP/DSI/0.0001
                 TYPE *, 'S, I, STEMP (mics), DPLD: ', SDUM, I, STEMPDUM, DUM
                IF (M.LT.TBINS-1) THEN
                    M=M+1
                    LTEMP (M) = ENERGY/STEMP
                    XVAL=LTEMP (M)
                    CALL INTERPOLATE INTLET (XVAL, N, L, FLUX, YVAL)
                    FTEMP (M) = YVAL
                ENDIF
  45
             CONTINUE
  50
        CONTINUE
C
C
        We now have the appropriate array of LET and FLUX values.
С
        For the numerical integration, these must be ordered.
C
        Use the INDEXX routine from Numerical Recipes:
        CALL INDEXX (M, TBINS, LTEMP, INDX)
```

```
RETURN
END
```

```
SUBROUTINE TEMP_STORAGE(NBINS, NVAL, LVAL, FVAL, N, L, FLUX)
        IMPLICIT NONE
        INTEGER*4 NBINS, NVAL, N, K
        REAL*4 LVAL, FVAL, L, FLUX
        DIMENSION LVAL(1), FVAL(1), L(1), FLUX(1)
С
        Copy Integral LET spectrum:
        DO 5 K=1, NVAL
                 L(K) = LVAL(K)
                 FLUX(K)=FVAL(K)
 5
        CONTINUE
        N=NVAL
        IF (N.GT. NBINS) THEN
        WRITE(6,9999) N, NBINS
 9999
        FORMAT('@ 10002 ABNORMAL TERMINATION: ',
           /,1x,' LET spec array out of bounds in GET_UPSET: ',
     &
          /,1x,'N = ',18,'NBINS = ',18,
          /,1x,' STOP.')
     &
        STOP
        ENDIF
        RETURN
        END
SUBROUTINE EXPAND_SAMPLING(NBINS, N, L, FLUX,
                                   LMIN, ENERGY, PMAX, X, Y, Z,
     &
                                   M, LUSE, FUSE)
        IMPLICIT NONE
        INTEGER*4 NBINS, N, M, TBINS
        REAL*4 L, FLUX, LMIN, ENERGY, PMAX, X, Y, Z, LUSE, FUSE
        DIMENSION L(1),FLUX(1),LUSE(1),FUSE(1)
        PARAMETER (TBINS=5000)
        REAL*4 LTEMP, FTEMP
        DIMENSION LTEMP (TBINS), FTEMP (TBINS)
        INTEGER*4 I, K
        REAL*4 S, STEMP, SDUM, XVAL, YVAL
        REAL*4 SCALE, SMALLEST, SAMPLE SIZE
        DATA SCALE/0.01/
        REAL*4 DSI
        DATA DSI/2.321/
C
       For idiot checks:
C
        REAL*4 DUM, DIFPLD, STEMPDUM
        INTEGER*4 INDX
        DIMENSION INDX (TBINS)
```

```
NOTE: X,Y,Z assumed here to be in g/cm2!!!
С
С
        Store relevant portion of input LET spectrum:
C
        DO 4 I=1,N
           IF (L(I).LT.LMIN) GO TO 4
           IF (FLUX(I).LE.0.0) GOTO 6
           M=M+1
           LTEMP(M) = L(I)
           FTEMP (M) = FLUX (I)
        CONTINUE
 4
        CONTINUE
 6
        Now we wish to do additional samplings around the peaks
C
        in the pathlength distribution.
С
        Specifically, the additional sampling is done on a scale
C
        equal to 1% of the smallest dimension, but no larger than 0.01
С
        microns. The sampling is done at 100 points ranging from
С
        x-10*scale to x+90*scale.
С
        PMAX=SQRT(X*X+Y*Y+Z*Z)
        SMALLEST=MIN(X,Y,Z)/0.0001/DSI
        SAMPLE_SIZE=SCALE*SMALLEST
        IF (SAMPLE_SIZE.GT.0.01) SAMPLE_SIZE=0.01
        DO 50 K=1,3
              IF (K.EQ.1) S=X
              IF (K.EQ.2) S=Y
              IF (K.EQ.3) S=Z
              Suppress redundant samplings:
С
              IF (K.EQ.2 .and. (ABS(X-Y).LE.0.0001*X)) GOTO 50
              IF (K.EQ.3 .and.
               (ABS(X-Z).LE.0.0001*X .or. ABS(Y-Z).LE.0.0001*Y)) GOTO 50
     &
              DO 45 I=-11,89
                 STEMP=S+FLOAT(I) *SAMPLE_SIZE*0.0001*DSI
                 IF (STEMP.LE.O. .or. STEMP.GT.PMAX) GOTO 45
        Idiot checks again:
\mathsf{C}
                  DUM=DIFPLD(STEMP, X, Y, Z)
С
                  SDUM=S/DSI/0.0001
C
                  STEMPDUM=STEMP/DSI/0.0001
C
                  TYPE *,' S,I,STEMP(mics),DPLD: ',SDUM,I,STEMPDUM,DUM
C
                  IF (M.LT.TBINS-1) THEN
                     M=M+1
                     LTEMP (M) = ENERGY / STEMP
                      XVAL=LTEMP(M)
                      CALL INTERPOLATE INTLET (XVAL, N, L, FLUX, YVAL)
                      FTEMP (M) =YVAL
                  ENDIF
               CONTINUE
   45
         CONTINUE
   50
 C
         We now have the appropriate array of LET and FLUX values.
 C
         For the numerical integration, these must be ordered.
 C
         Use the INDEXX routine from Numerical Recipes:
 C
         CALL INDEXX (M, TBINS, LTEMP, INDX)
```

```
C
        Now store the values according to increasing LET:
        DO 55 I=1, M
              IF (I.LE.NBINS.AND.I.LE.TBINS
                             .AND.INDX(I).LE.TBINS) THEN
     &
                  LUSE (I) = LTEMP (INDX (I))
                  FUSE(I) = FTEMP(INDX(I))
              ELSE
                  WRITE(6,9999) M, NBINS, TBINS, I, INDX(I)
                  FORMAT(' FATAL ERROR IN GET_UPSET:',
9999
                       /,'
                                M, NBINS, TBINS: ',316,
                       /,'
                                 I, INDX(I): ',216)
     &
              ENDIF
  55
        CONTINUE
        RETURN
        END
        SUBROUTINE INTERPOLATE INTLET(X,N,L,FLUX,Y)
C
C
        Does a linear interpolation on a log-log plot of the
C
        integral flux vs. LET curve.
C
        IMPLICIT NONE
        INTEGER*4 N
        REAL*4 X,Y,L,FLUX
        DIMENSION L(1), FLUX(1)
        REAL*8 X1, X2, X3, Y1, Y2, Y3, SLOPE
        INTEGER*4 I
        IF (X.LE.L(1)) THEN
            Y=FLUX(1)
        ELSEIF (X.GT.L(N)) THEN
            Assume integral flux vanishes above the highest L value.
            Y=0.0
        ELSE
            DO 100 I=1, N-1
               IF (X.LE.L(I+1)) THEN
                   IF (FLUX(I).GT.0. .and. FLUX(I+1).GT.0.) THEN
                       X1=ALOG(L(I))
                       Y1=ALOG(FLUX(I))
                       X2=ALOG(L(I+1))
                       Y2=ALOG(FLUX(I+1))
                       SLOPE = (Y2-Y1)/(X2-X1)
                       X3 = ALOG(X)
                       Y3=SLOPE*(X3-X1)+Y1
                       Y=EXP(Y3)
                       GOTO 150
                   ELSE
                        Y=0.0
                   ENDIF
               ENDIF
100
            CONTINUE
150
            CONTINUE
        ENDIF
        RETURN
        END
```

C C

C

C

C

C

This version uses the Epoch 1980.0 vertical cutoff grid of Shea and Smart.

IMPLICIT NONE

INTEGER I,J
REAL CUTOFF(33,72),CN,CS

COMMON/CUTOFF80/CUTOFF, CN, CS

This common block contains the table of world wide vertical geomagnetic cutoffs at 20 km altitude, tabulated every 5 degrees in latitude (to +/- 80 degrees) and 5 degrees in longitude. It was given to JHA by private communication from D.F. Smart on 11/27/89. This calculation is for Epoch 1980.0, ie. the cutoff calculation used the 10th degree IGRF model (1980), as discussed in Shea & Smart, Proc. 18th ICRC, v. 3, p. 415 (1983).

```
DATA (CUTOFF(1, J), J=1,72)/
#
           0.31, 0.29, 0.26, 0.23, 0.21, 0.18, 0.16, 0.14,
     0.34,
#
     0.12,
           0.10, 0.09, 0.07, 0.05,
                                    0.04, 0.02,
                                                 0.01, 0.01,
#
     0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00,
#
     0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.01, 0.01,
#
     0.02, 0.04, 0.05, 0.07,
                              0.09, 0.10, 0.12, 0.15, 0.17,
#
     0.20, 0.23, 0.25, 0.28,
                              0.32, 0.35, 0.39,
                                                 0.41,
#
     0.45, 0.47, 0.50, 0.52, 0.53, 0.53, 0.54,
                                                 0.54, 0.53,
#
     0.53, 0.52,
                  0.51, 0.50, 0.46,
                                    0.43, 0.39,
                                                 0.37, 0.36/
   DATA (CUTOFF(2,J),J=1,72)/
#
     0.59, 0.51, 0.45, 0.42, 0.37,
                                    0.34, 0.28,
                                                 0.25, 0.21,
#
     0.18,
           0.15, 0.12, 0.10, 0.08,
                                    0.06, 0.04,
                                                 0.02, 0.00,
#
     0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00,
     0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.02,
#
     0.04, 0.07, 0.09, 0.11, 0.15, 0.18, 0.22,
#
                                                 0.24, 0.30,
     0.34, 0.42, 0.48, 0.53, 0.60, 0.65, 0.70,
#
                                                 0.75, 0.83,
#
     0.87, 0.91, 0.90, 1.01, 1.00, 1.07, 1.04,
                                                 1.03, 1.03,
#
     0.98, 0.93, 0.93, 0.86, 0.82, 0.79, 0.72,
                                                 0.66, 0.60/
   DATA (CUTOFF(3,J),J=1,72)/
#
     0.84, 0.79, 0.73, 0.66, 0.58, 0.50, 0.46,
                                                 0.39, 0.36,
#
     0.30,
          0.24, 0.21, 0.16, 0.13, 0.10, 0.08,
                                                 0.05, 0.03,
#
     0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00,
                                                 0.00, 0.00,
#
     0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.02,
                                                 0.05, 0.08,
     0.10, 0.14, 0.18, 0.22, 0.27, 0.30, 0.40,
#
                                                 0.47, 0.53,
#
     0.60,
           0.72, 0.80, 0.90, 1.06, 1.11, 1.20,
                                                 1.30, 1.43,
#
     1.51, 1.61, 1.64, 1.67, 1.72, 1.74, 1.75,
                                                 1.70, 1.65,
#
     1.59, 1.52, 1.42, 1.33, 1.27,
                                    1.22, 1.11,
                                                 1.04, 0.95/
   DATA (CUTOFF(4,J),J=1,72)/
#
     1.29, 1.15, 1.03, 0.96, 0.85,
                                    0.76, 0.72,
                                                 0.60, 0.57,
#
     0.48,
          0.41, 0.35, 0.29, 0.25,
                                    0.19, 0.14,
                                                 0.13, 0.09,
#
     0.06, 0.04, 0.02, 0.00, 0.00, 0.00, 0.00,
                                                 0.00,
#
     0.00, 0.00, 0.02, 0.03, 0.05, 0.08, 0.10,
                                                 0.14,
                                                        0.18,
     0.22, 0.27, 0.35, 0.42, 0.49, 0.61, 0.69,
#
                                                 0.80,
#
           1.21, 1.34, 1.55, 1.65,
     1.05,
                                    1.86, 1.96,
                                                 2.14, 2.32,
#
     2.46, 2.53, 2.67, 2.72,
                              2.70, 2.71, 2.63, 2.56, 2.51,
     2.40, 2.27, 2.20, 2.02, 1.87,
                                    1.70, 1.61,
                                                 1.45,
                                                       1.35/
   DATA (CUTOFF(5,J),J=1,72)/
#
     1.69, 1.53, 1.45, 1.29, 1.15, 1.11, 1.03, 0.94,
                                                      0.83.
#
     0.76, 0.65, 0.59, 0.49, 0.42, 0.36, 0.30, 0.25, 0.20,
```

```
0.15, 0.13, 0.11, 0.09, 0.08, 0.07, 0.06, 0.06, 0.07,
     0.08, 0.09, 0.12, 0.14, 0.16, 0.22, 0.26,
                                                 0.32,
                                                        0.39,
#
                                     1.07, 1.20, 1.38, 1.54,
           0.58, 0.68, 0.76, 0.93,
     0.51,
                                           2.98,
           1.93, 2.10, 2.27, 2.64,
                                                 3.23,
                                                        3.49,
                                     2.76,
     1.77,
           3.79, 3.91, 4.01, 3.94, 3.94, 3.82,
                                                 3.71,
                                                        3.57,
     3.75,
                                     2.35, 2.14,
                                                 2.01,
                                                        1.78/
           3.21, 2.98, 2.72, 2.52,
#
     3.39,
   DATA (CUTOFF(6,J),J=1,72)/
     2.17, 1.97, 1.86, 1.72, 1.66, 1.56, 1.45,
                                                 1.30, 1.20,
#
     1.14, 1.00, 0.92, 0.82, 0.73, 0.65, 0.56, 0.47, 0.41,
#
           0.31, 0.27, 0.24, 0.22, 0.21, 0.21, 0.21,
#
     0.35,
     0.25, 0.28, 0.32, 0.38, 0.43,
                                    0.51, 0.59, 0.67,
                                                        0.80,
                                                 2.18,
     0.95, 1.07, 1.22, 1.36, 1.58, 1.76, 1.94,
                                                        2.33,
     2.64, 2.80, 3.03, 3.35, 3.80, 4.02, 4.29, 4.59, 4.75,
#
     4.90, 5.07, 5.10, 5.18, 5.17, 5.00, 4.90, 4.74,
                                                        4.54,
     4.25, 3.98, 3.70, 3.48, 3.21,
                                    2.98, 2.77,
                                                  2.51,
                                                        2.32/
#
   DATA (CUTOFF(7,J),J=1,72)/
     2.76, 2.55, 2.38, 2.27, 2.09, 2.02, 1.97,
                                                 1.85, 1.73,
#
     1.64, 1.52, 1.42, 1.30, 1.19, 1.05, 0.94, 0.86, 0.78,
#
     0.71, 0.67, 0.58, 0.56, 0.53, 0.53, 0.53, 0.51, 0.56,
#
           0.66, 0.71, 0.81, 0.89, 1.01, 1.15, 1.27, 1.42,
     0.60,
#
           1.83, 2.03, 2.24, 2.51, 2.70, 2.94, 3.15, 3.40,
#
     1.65,
           4.05, 4.20, 4.58, 4.85,
                                     5.29, 5.61, 6.03, 6.62,
#
   . 3.79,
                                            6.77, 6.37, 5.95,
     7.05, 7.36, 7.42, 7.41, 7.32,
                                     7.10,
#
                                                        2.87/
           5.01, 4.60, 4.27, 4.00,
                                     3.62, 3.42,
                                                 3.14,
     5.47,
#
   DATA (CUTOFF(8,J),J=1,72)/
     3.33, 3.22, 3.01, 2.88, 2.80, 2.67, 2.53,
                                                 2.50, 2.47,
#
     2.38, 2.29, 2.10, 2.00, 1.82, 1.69, 1.54, 1.47, 1.35,
#
           1.18, 1.14, 1.12, 1.08, 1.08, 1.10, 1.13, 1.16,
#
     1.28,
           1.33, 1.36, 1.47, 1.67, 1.79,
                                            2.01, 2.17, 2.43,
#
     1.22,
     2.69, 2.96, 3.15, 3.30, 3.70,
                                     4.08, 4.28,
                                                 4.42,
                                                        4.63,
#
     4.99, 5.29, 5.61, 6.01, 6.63, 7.41,
                                            7.87, 8.16, 8.69,
#
     8.89, 8.97, 8.77, 8.52, 8.31, 8.01, 7.74, 7.58, 7.16,
#
            6.43, 5.89, 5.34, 4.85, 4.43,
                                           4.15,
                                                 3.82,
                                                        3.52/
#
     6.80,
   DATA (CUTOFF(9,J),J=1,72)/
     4.03, 3.85, 3.72, 3.62, 3.49, 3.41, 3.38,
                                                 3.33, 3.33,
#
     3.26, 3.18, 3.00, 2.89, 2.77, 2.69, 2.58, 2.35, 2.25,
#
                                            2.04, 2.01,
                                                        2.14,
     2.21, 2.06, 2.02, 2.04, 2.05, 2.05,
#
     2.22, 2.31, 2.43, 2.53, 2.86, 2.96, 3.29, 3.48, 3.74,
#
            4.48, 4.76, 4.84, 5.05, 5.30, 5.60, 5.96, 6.31,
     4.15,
#
                  7.91, 8.18, 8.55, 9.11, 9.73, 10.05, 10.14,
           7.27,
#
     6.76,
           9.98, 9.81, 9.63, 9.39, 9.13,
                                           8.80, 8.32, 7.88,
#
     10.08,
                                            5.00,
                                                 4.60,
                                                        4.34/
                               5.94, 5.48,
      7.35, 6.82, 6.68, 6.18,
#
    DATA (CUTOFF(10,J),J=1,72)/
      5.05, 4.79, 4.65, 4.49, 4.42, 4.39, 4.37, 4.33, 4.39,
#
            4.34, 4.26, 4.12, 4.12, 3.99,
                                           3.85, 3.68,
                                                         3.52,
#
      4.38,
                                                        3.45,
      3.47, 3.40, 3.40, 3.41, 3.35, 3.35, 3.34, 3.37,
#
      3.55, 3.69, 3.89, 4.10, 4.35, 4.51, 4.90, 5.18, 5.38,
#
           5.95, 6.25, 6.62, 7.01, 7.47, 8.01, 8.51, 8.84,
#
      5.65,
      8.73, 8.41, 9.00, 9.62, 10.73, 10.95, 11.05, 11.10, 11.12,
#
     11.09, 10.95, 10.75, 10.55, 10.29, 10.01, 9.73, 9.39, 9.00,
#
      8.50, 7.99, 7.47, 6.88, 6.52, 6.12,
                                            5.95, 5.66,
                                                         5.32/
#
    DATA (CUTOFF(11, J), J=1, 72)/
                                            5.84,
                                                  5.82,
                                                         5.91,
                              5.85, 5.83,
      6.11, 5.99, 5.86, 5.84,
#
                                                  5.27, 5.23,
      5.80, 5.74, 5.72, 5.58, 5.51, 5.45,
                                            5.40,
#
      5.19, 5.14, 5.18, 5.14, 5.13, 5.06,
                                            5.09, 5.21, 5.22,
#
      5.38, 5.45, 5.60, 5.84, 6.02, 6.32, 6.60, 6.91, 7.46,
#
      7.98, 8.58, 9.05, 9.45, 9.84, 9.25,
                                           9.40,
                                                  9.59, 10.12,
#
     10.75, 11.25, 11.69, 11.79, 11.86, 11.91, 11.92, 11.91, 11.87,
     11.78, 11.67, 11.53, 11.34, 11.12, 10.85, 10.56, 10.29,
#
      9.57, 9.15, 8.65, 8.15, 7.62, 7.14, 6.88, 6.74, 6.33/
```

```
DATA (CUTOFF (12, J), J=1,72)/
                                 7.45, 7.52,
                                                       7.73,
                                                              7.82,
            7.32, 7.35, 7.29,
                                                7.63,
                                                              7.37,
            8.02, 8.03, 8.02, 8.02, 7.88,
                                               7.71,
                                                       7.53,
#
      7.93,
                           7.26, 7.35, 7.35, 7.42,
                                                       7.44,
                                                               7.55,
                    7.30,
             7.29,
#
      7.28,
            7.82, 8.06, 8.36, 8.75, 9.24, 9.49, 9.96, 10.40,
     10.19, 10.44, 10.76, 11.24, 11.61, 11.82, 11.98, 12.26, 12.48,
#
     12.56, 12.62, 12.65, 12.67, 12.68, 12.66, 12.63, 12.57, 12.49,
#
     12.38, 12.25, 12.10, 11.94, 11.76, 11.59, 11.40, 11.13, 10.84,
#
     10.52, 10.21, 9.79, 9.35, 8.95, 8.50, 8.04, 7.78,
#
    DATA (CUTOFF(13,J),J=1,72)/
      8.87, 8.81, 8.79, 8.89, 8.99, 9.05, 9.26, 9.42,
                                                               9.60,
#
      9.74, 9.92, 10.02, 10.24, 10.42, 10.26, 10.45, 10.52, 10.53,
#
     10.55, 10.58, 10.64, 10.69, 10.73, 10.69, 10.75, 10.56, 10.63,
#
     10.66, 10.38, 10.49, 10.62, 10.90, 11.30, 11.87, 12.27, 12.56,
#
     12.72, 12.86, 13.02, 13.15, 13.23, 13.31, 13.36, 13.40, 13.41,
     13.42, 13.41, 13.38, 13.35, 13.30, 13.25, 13.17, 13.08, 12.98,
     12.85, 12.71, 12.57, 12.42, 12.28, 12.14, 11.99, 11.85, 11.69,
#
     11.47, 11.28, 10.84, 10.49, 10.14, 9.75, 9.47, 9.19,
#
    DATA (CUTOFF(14,J),J=1,72)/
     10.45, 10.49, 10.61, 10.63, 10.75, 10.92, 11.08, 11.32, 11.57,
#
     11.75, 11.92, 12.13, 12.32, 12.48, 12.70, 12.80, 12.91, 12.99,
#
     13.06, 13.14, 13.20, 13.24, 13.23, 13.20, 13.17, 13.13, 13.13,
#
     13.18, 13.25, 13.32, 13.39, 13.48, 13.58, 13.69, 13.81, 13.93,
#
     14.01, 14.09, 14.14, 14.17, 14.17, 14.17, 14.15, 14.12, 14.08,
#
     14.03, 13.98, 13.91, 13.84, 13.76, 13.67, 13.57, 13.46, 13.33,
     13.19, 13.04, 12.90, 12.77, 12.65, 12.56, 12.47, 12.38, 12.28,
     12.15, 11.98, 11.78, 11.53, 11.32, 11.08, 10.79, 10.62, 10.50/
#
    DATA (CUTOFF(15, J), J=1,72)/
     11.86, 11.92, 12.01, 12.11, 12.23, 12.39, 12.57, 12.77, 12.99,
#
     13.23, 13.46, 13.69, 13.90, 14.09, 14.26, 14.40, 14.52, 14.62,
#
     14.71, 14.79, 14.85, 14.88, 14.89, 14.87, 14.84, 14.79, 14.73,
#
     14.69, 14.66, 14.65, 14.65, 14.68, 14.72, 14.77, 14.81, 14.86,
#
     14.87, 14.87, 14.86, 14.82, 14.78, 14.72, 14.66, 14.59, 14.51,
#
     14.43, 14.35, 14.25, 14.15, 14.05, 13.93, 13.81, 13.67, 13.52,
#
     13.37, 13.22, 13.08, 12.97, 12.89, 12.84, 12.81, 12.78, 12.75,
#
     12.69, 12.60, 12.49, 12.36, 12.21, 12.05, 11.92, 11.85, 11.85/
#
    DATA (CUTOFF(16, J), J=1,72)/
     12.94, 13.03, 13.14, 13.27, 13.42, 13.58, 13.77, 13.97, 14.18,
     14.41, 14.64, 14.88, 15.10, 15.30, 15.48, 15.64, 15.77, 15.88,
#
     15.97, 16.03, 16.07, 16.07, 16.05, 16.01, 15.94, 15.86, 15.77,
 #
      15.68, 15.61, 15.54, 15.50, 15.46, 15.44, 15.42, 15.41, 15.39,
 #
     15.35, 15.31, 15.25, 15.17, 15.09, 15.01, 14.92, 14.82, 14.72,
      14.62, 14.51, 14.40, 14.28, 14.15, 14.01, 13.86, 13.70, 13.53,
 #
      13.36, 13.21, 13.08, 12.99, 12.95, 12.95, 12.98, 13.03, 13.07,
 #
      13.09, 13.08, 13.05, 13.00, 12.93, 12.88, 12.85, 12.84, 12.87/
 #
     DATA (CUTOFF(17, J), J=1,72)/
      13.80, 13.93, 14.07, 14.22, 14.38, 14.55, 14.73, 14.92, 15.12,
 #
      15.34, 15.56, 15.78, 16.00, 16.21, 16.39, 16.56, 16.69, 16.79,
 #
      16.87, 16.91, 16.92, 16.90, 16.85, 16.77, 16.67, 16.56, 16.43,
 #
      16.31, 16.18, 16.07, 15.97, 15.89, 15.81, 15.74, 15.67, 15.60,
      15.53, 15.44, 15.35, 15.25, 15.15, 15.04, 14.93, 14.83, 14.71,
 #
      14.60, 14.48, 14.35, 14.21, 14.06, 13.89, 13.71, 13.52, 13.32,
 #
      13.14, 12.98, 12.86, 12.80, 12.80, 12.87, 12.97, 13.09, 13.21,
 #
      13.32, 13.40, 13.45, 13.48, 13.50, 13.52, 13.55, 13.61, 13.69/
     DATA (CUTOFF(18,J),J=1,72)/
      14.41, 14.56, 14.72, 14.88, 15.05, 15.21, 15.38, 15.56, 15.74,
      15.94, 16.14, 16.36, 16.57, 16.77, 16.95, 17.11, 17.25, 17.34,
 #
      17.41, 17.43, 17.42, 17.38, 17.30, 17.20, 17.07, 16.93, 16.77,
 #
      16.61, 16.45, 16.30, 16.15, 16.01, 15.89, 15.77, 15.66, 15.55,
 #
      15.43, 15.31, 15.19, 15.07, 14.96, 14.84, 14.73, 14.61, 14.49,
```

```
#
    14.36, 14.22, 14.08, 13.92, 13.72, 13.51, 13.22, 12.99, 12.71,
    12.45, 12.34, 12.34, 12.29, 12.41, 12.55, 12.73, 12.93, 13.14,
#
#
     13.33, 13.50, 13.64, 13.76, 13.85, 13.94, 14.04, 14.14, 14.27/
   DATA (CUTOFF(19,J),J=1,72)/
     14.70, 14.86, 15.02, 15.19, 15.35, 15.51, 15.67, 15.82, 15.99,
#
    16.17, 16.35, 16.55, 16.75, 16.95, 17.13, 17.29, 17.42, 17.52,
#
#
    17.57, 17.59, 17.56, 17.50, 17.41, 17.29, 17.15, 16.99, 16.81,
    16.63, 16.44, 16.25, 16.06, 15.89, 15.72, 15.56, 15.41, 15.25,
#
    15.10, 14.96, 14.81, 14.68, 14.55, 14.42, 14.30, 14.18, 14.03,
#
    13.88, 13.74, 13.56, 13.35, 13.13, 12.75, 12.36, 12.00, 11.59,
#
    11.27, 11.05, 10.96, 11.06, 11.41, 11.85, 12.22, 12.53, 12.83,
#
    13.11, 13.36, 13.58, 13.78, 13.95, 14.10, 14.25, 14.39, 14.54/
#
   DATA (CUTOFF(20,J), J=1,72)/
    14.63, 14.79, 14.95, 15.10, 15.25, 15.40, 15.54, 15.68, 15.83,
    15.99, 16.16, 16.34, 16.54, 16.73, 16.91, 17.07, 17.20, 17.29,
#
    17.35, 17.36, 17.34, 17.27, 17.18, 17.06, 16.91, 16.75, 16.56,
#
    16.37, 16.16, 15.95, 15.75, 15.54, 15.34, 15.14, 14.95, 14.77,
#
    14.59, 14.41, 14.25, 14.09, 13.95, 13.81, 13.67, 13.52, 13.35,
#
#
    13.19, 12.99, 12.77, 12.46, 12.06, 11.60, 11.05, 10.41, 9.86,
     9.27, 8.69, 8.41, 8.64, 9.15, 10.07, 11.01, 11.75, 12.22,
#
    12.60, 12.94, 13.24, 13.50, 13.73, 13.94, 14.12, 14.30, 14.46/
#
   DATA (CUTOFF(21,J),J=1,72)/
    14.17, 14.32, 14.47, 14.61, 14.74, 14.87, 14.99, 15.12, 15.25,
#
    15.39, 15.55, 15.72, 15.91, 16.09, 16.27, 16.43, 16.57, 16.67,
#
    16.73, 16.75, 16.73, 16.68, 16.60, 16.49, 16.36, 16.21, 16.03,
#
    15.85, 15.64, 15.43, 15.21, 14.99, 14.76, 14.54, 14.32, 14.11,
#
#
    13.90, 13.70, 13.51, 13.33, 13.16, 13.00, 12.84, 12.64, 12.45,
    12.23, 11.96, 11.67, 11.32, 10.89, 10.28, 9.35, 8.60, 8.31,
#
#
     7.67, 7.15, 6.79, 6.61, 7.24, 8.16, 8.49, 9.68, 10.89,
    11.67, 12.18, 12.55, 12.88, 13.17, 13.42, 13.63, 13.83, 14.00/
#
   DATA (CUTOFF(22,J),J=1,72)/
    13.25, 13.38, 13.53, 13.68, 13.81, 13.92, 14.03, 14.14, 14.25,
#
#
    14.38, 14.52, 14.68, 14.86, 15.04, 15.22, 15.37, 15.52, 15.62,
    15.69, 15.73, 15.73, 15.70, 15.65, 15.57, 15.47, 15.36, 15.22,
#
    15.05, 14.87, 14.67, 14.46, 14.23, 14.00, 13.76, 13.52, 13.28,
#
    13.04, 12.81, 12.58, 12.38, 12.17, 11.96, 11.74, 11.49, 11.21,
#
    10.88, 10.53, 10.10, 9.63, 9.06, 8.44, 7.74, 7.09, 6.49,
#
#
      6.02, 5.69, 5.59, 5.51, 5.78, 6.15, 7.02, 8.04, 9.04,
      9.98, 10.81, 11.34, 11.78, 12.14, 12.42, 12.68, 12.90, 13.07/
#
   DATA (CUTOFF(23,J),J=1,72)/
#
    11.49, 11.62, 11.74, 11.83, 11.91, 11.98, 12.10, 12.19, 12.35,
    12.51, 12.67, 12.83, 13.09, 13.34, 13.59, 13.82, 14.00, 14.11,
#
    14.19, 14.27, 14.30, 14.31, 14.31, 14.28, 14.23, 14.17, 14.08,
#
#
    13.97, 13.82, 13.66, 13.46, 13.25, 13.02, 12.75, 12.43, 12.07,
#
    11.69, 11.26, 10.79, 10.48, 10.14, 9.80, 9.65, 9.41, 9.10,
#
      8.78, 8.25, 7.62, 6.99, 6.40, 5.97, 5.54, 5.08, 4.74,
#
      4.38,
             4.19, 4.10, 4.02, 4.21, 4.61, 5.14, 5.65, 6.46,
#
            8.64, 9.38, 10.02, 10.52, 10.72, 10.94, 11.16, 11.33/
   DATA (CUTOFF(24,J),J=1,72)/
#
      9.77, 9.75, 9.86, 9.74, 9.76, 9.84, 10.01, 10.17, 10.27,
#
    10.42, 10.45, 10.72, 10.88, 11.20, 11.37, 11.27, 11.33, 11.34,
    11.39, 11.46, 11.56, 11.63, 11.74, 11.87, 11.95, 12.08, 12.17,
#
    12.18, 12.09, 11.92, 11.69, 11.37, 11.04, 10.67, 10.31,
#
#
      9.52, 9.44, 9.49, 8.99, 8.52, 8.15, 7.67, 7.07,
#
      6.11, 5.75, 5.48, 5.17, 4.89, 4.56, 4.21, 3.80,
#
      3.24,
            3.08, 2.98, 2.99, 3.13, 3.41, 3.83,
                                                      4.15,
                                                             4.74,
      5.26, 5.76, 6.45, 7.36,
                                        8.72, 9.11,
#
                                 8.14,
                                                      9.38,
   DATA (CUTOFF(25, J), J=1, 72)/
#
      6.95, 7.19, 7.34, 7.44,
                                 7.50,
                                        7.52, 7.59, 7.58,
                                                             7.63,
            7.83, 7.91, 8.07, 8.24, 8.38, 8.54, 8.71,
#
      7.73,
```

```
9.46, 9.57,
                                                   9.79, 9.90,
     8.99, 9.03, 9.13, 9.23, 9.30,
#
                                                   8.64,
                                                          8.31,
     9.99, 10.01, 9.99, 9.82, 9.63,
                                      9.37, 9.05,
#
                                           5.49,
                                                   5.27,
                                                          4.89,
     7.96, 7.41, 6.90, 6.47, 6.08,
                                      5.78,
     4.60, 4.34, 4.02, 3.63, 3.30,
                                      3.08,
                                            2.77,
                                                   2.54,
                                                          2.30,
                                      2.27, 2.57,
                                                   2.86,
                                                          3.22,
     2.14, 2.09, 2.04, 2.05, 2.13,
                                            6.15,
                                                   6.42,
                                                          6.72/
     3.79, 4.18, 4.63, 5.08, 5.41,
                                      5.76,
   DATA (CUTOFF(26,J),J=1,72)/
                                      5.34, 5.35,
                                                          5.32,
                                                   5.36,
     4.99, 5.02, 5.09, 5.20, 5.21,
                                      5.73, 5.81,
                                                   5,98, 6.03,
           5.52, 5.53, 5.66, 5.71,
     5.44,
#
                                                   6.75, 6.85,
     6.08, 6.16, 6.19, 6.31, 6.37,
                                     6.45, 6.59,
#
                                      6.66, 6.38,
                                                   6.17,
                                                          5.83,
           7.03, 6.99, 6.96, 6.88,
#
     6.96,
                                      4.40, 4.15,
                                                   3.77,
                                                          3.47,
     5.60, 5.35, 5.03, 4.85, 4.59,
#
                                      1.88, 1.75, 1.58,
     3.16, 2.92, 2.66, 2.37, 2.10,
#
                                      1.47, 1.62,
     1.35, 1.28, 1.25, 1.27, 1.36,
                                                   1.87,
                                                          2.11,
#
                                                   4.61,
                                                          4.68/
     2.47, 2.71, 3.06, 3.35, 3.80,
                                      4.17,
                                            4.35,
#
   DATA (CUTOFF(27, J), J=1,72)/
                                      3.76, 3.83,
                                                   3.86,
                                                          3.91,
     3.32, 3.35, 3.49, 3.59, 3.69,
                                                          4.32,
     3.94, 3.97, 4.03, 4.06, 4.11,
                                      4.17,
                                            4.20,
                                                   4.32,
#
                                      4.64, 4.69,
                                                   4.85,
                                                         5.00,
     4.34, 4.36, 4.44, 4.45, 4.49,
#
            5.05, 4.98, 4.97, 4.86, 4.77, 4.69,
                                                   4.54, 4.47,
#
     5.00,
                                      3.00,
                                                   2.51,
                                                          2.25,
            4.08, 3.78, 3.46, 3.21,
                                             2.81,
     4.25,
                                             0.98,
                                                   0.87,
                                                          0.81,
      2.05, 1.79, 1.64, 1.42, 1.28,
                                      1.11,
#
                                                          1.27,
      0.75, 0.71, 0.72, 0.73, 0.79, 0.87,
                                             0.95,
                                                   1.10,
#
                         2.13, 2.33, 2.65,
                                             2.85,
                                                   2.96,
                                                          3.18/
            1.67, 1.92,
#
      1.43,
    DATA (CUTOFF(28,J),J=1,72)/
                                                   2.47,
            2.08, 2.26, 2.29, 2.36, 2.41, 2.45,
                                                          2.51,
#
      2.00,
            2.64, 2.64, 2.67, 2.68, 2.69, 2.73,
                                                   2.77,
                                                          2.78,
#
      2.53,
           2.80, 2.86, 2.93, 2.93, 3.03,
                                             3.12, 3.19,
                                                          3.26,
#
      2.84,
      3.31, 3.38, 3.38, 3.31, 3.30, 3.26,
                                             3.15,
                                                   3.08,
                                                          3.02,
#
      2.85, 2.69, 2.49, 2.28, 2.18, 1.96, 1.75, 1.55,
                                                          1.37,
#
            1.07, 0.90, 0.81, 0.70, 0.58, 0.49, 0.44,
                                                          0.41,
      1.23,
#
            0.36, 0.35, 0.36, 0.38, 0.46, 0.50, 0.58,
                                                          0.68,
      0.36,
#
                                      1.54, 1.70,
                                                   1.81,
                                                          1.96/
            0.93, 1.05, 1.21, 1.35,
#
      0.80,
    DATA (CUTOFF(29,J),J=1,72)/
      1.16, 1.23, 1.29, 1.36, 1.39, 1.44, 1.43,
                                                   1.52,
                                                          1.54,
#
            1.60, 1.57, 1.62, 1.64, 1.69, 1.68, 1.67,
                                                          1.69,
#
      1.59,
      1.70, 1.74, 1.73, 1.80, 1.86, 1.89, 1.96, 1.99,
                                                          2.06,
#
      2.05, 2.09, 2.09, 2.12, 2.14, 2.04, 2.06,
                                                   1.96,
                                                          1.86,
#
            1.63, 1.55, 1.38, 1.27, 1.16, 0.98,
                                                   0.89,
                                                          0.77,
      1.76,
#
            0.56, 0.46, 0.40,
                                             0.23, 0.20,
                                                          0.17,
                               0.32,
                                     0.27,
#
      0.65,
                                                   0.27,
                                                          0.34,
                                      0.21,
                                             0.23,
            0.15, 0.15, 0.16,
                                0.17,
#
      0.16,
                                             0.90,
                                                   1.00,
                                                          1.09/
                                0.72,
                                      0.85,
#
      0.39,
            0.44,
                   0.53,
                         0.64,
    DATA (CUTOFF(30, J), J=1,72)/
      0.60, 0.62, 0.66, 0.69, 0.76,
                                     0.77,
                                             0.80,
                                                   0.82,
                                                          0.87,
#
            0.90, 0.88, 0.91, 0.93, 0.94,
                                             0.94,
                                                   0.98,
                                                          0.98,
#
      0.87,
      0.98, 1.00, 1.06, 1.03, 1.04, 1.09,
                                             1.12,
                                                   1.14,
                                                          1.16,
#
                                             1.11, 1.09,
                  1.22, 1.18, 1.18, 1.18,
                                                          1.02,
#
      1.23,
            1.14,
                   0.83, 0.77,
                                      0.57,
                                             0.49,
                                                   0.45,
                                                          0.39,
#
      0.99,
            0.90,
                                0.68,
                                             0.09,
                                                   0.07,
                                                          0.06,
            0.26,
                   0.21, 0.17,
                                0.13, 0.10,
#
      0.31,
                   0.05,
                         0.05,
                                0.06,
                                       0.07,
                                             0.09,
                                                    0.11,
                                                           0.14,
            0.05,
#
      0.05,
                                             0.43, 0.50,
                                                           0.54/
                                0.33,
                                      0.38,
                   0.25,
                         0.30,
#
      0.17,
            0.20,
    DATA (CUTOFF(31,J),J=1,72)/
                                             0.39,
                                                    0.42,
                                                           0.41,
            0.29, 0.31, 0.34,
                                      0.38,
#
      0.27,
                                0.36,
                                                    0.51,
                                                           0.50,
#
            0.45,
                   0.47, 0.48,
                                0.49,
                                      0.48,
                                             0.51,
      0.44,
                   0.56, 0.55,
                                0.55,
                                      0.56,
                                             0.58,
                                                   0.57,
                                                           0.58,
 #
      0.51,
            0.54,
                   0.61, 0.61,
                                0.59,
                                      0.57,
                                             0.57,
                                                   0.53,
                                                           0.51,
 #
            0.62,
      0.60,
                                             0.23, 0.19,
                                                           0.16,
                                0.30, 0.27,
                   0.38, 0.34,
 #
      0.47,
            0.45,
                   0.08, 0.06,
                                0.04, 0.02,
                                             0.00,
                                                    0.00,
                                                           0.00,
            0.10,
      0.13,
                                0.00, 0.00,
                                                    0.02,
                                             0.00,
                                                           0.04,
      0.00,
            0.00,
                   0.00, 0.00,
                                             0.19,
                                                    0.22,
                                                           0.25/
                   0.09, 0.11,
                                0.14,
                                      0.16,
      0.06,
            0.07,
```

C C

```
0.10, 0.11, 0.13, 0.14, 0.15, 0.16, 0.17, 0.19, 0.19,
                                                     0.24,
            0.21, 0.22, 0.22, 0.23, 0.24, 0.22,
                                                             0.24,
#
     0.20,
                                       0.28, 0.26, 0.27,
                                                             0.27,
           0.25, 0.25, 0.25, 0.26,
#
     0.25.
                                       0.25, 0.24,
            0.27, 0.27, 0.26, 0.25,
                                                     0.22,
                                                             0.20,
     0.26,
            0.18, 0.15, 0.14, 0.12,
                                       0.10,
                                              0.09,
                                                     0.07,
                                                             0.05,
     0.19,
            0.02, 0.00, 0.00, 0.00,
                                        0.00,
                                              0.00,
                                                      0.00,
                                                             0.00,
     0.03,
#
            0.00, 0.00, 0.00, 0.00,
                                        0.00,
                                               0.00,
                                                      0.00,
                                                             0.00,
     0.00,
     0.00, 0.00, 0.00, 0.02, 0.04,
                                        0.05,
                                               0.07,
                                                      0.08,
                                                             0.09/
#
   DATA (CUTOFF(33,J),J=1,72)/
     0.02, 0.03, 0.04, 0.05, 0.05,
                                               0.06,
                                                     0.07,
                                       0.06,
#
                                       0.09, 0.09, 0.09,
            0.08, 0.09, 0.09, 0.09,
                                                             0.10,
#
     0.08,
                                               0.11,
                                                     0.11,
                                                            0.10,
     0.10, 0.10, 0.10, 0.10, 0.10,
                                       0.11,
#
                                               0.08, 0.08, 0.07,
            0.10, 0.10, 0.10, 0.09, 0.09,
#
     0.10,
                          0.03, 0.02,
                                       0.01,
                                               0.00, 0.00,
            0.06, 0.04,
#
     0.07.
                                       0.00, 0.00,
                                                             0.00,
     0.00, 0.00, 0.00,
                          0.00, 0.00,
                                                     0.00,
                                       0.00, 0.00, 0.00,
     0.00, 0.00, 0.00, 0.00, 0.00,
                                                             0.00,
#
     0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.00, 0.01,
                                                             0.01/
   DATA CN,CS/ 0.05, 0.21/
    INIGRID=.TRUE.
   RETURN
   END
   SUBROUTINE InitPreCalcs (RigBins)
   IMPLICIT NONE
   INTEGER I, J, Nrigs
   PARAMETER (Nrigs=1001)
   REAL RigBins (Nrigs), RIGPC (Nrigs)
   REAL PCGTF1 (Nrigs), PCGTF2 (Nrigs), PCGTF3 (Nrigs), PCGTF4 (Nrigs)
   COMMON/PreCalcCMN/PCGTF1, PCGTF2, PCGTF3, PCGTF4
   DATA (RIGPC(I), I=1,90)/
     0.000, 0.020, 0.040, 0.060, 0.080, 0.100, 0.120, 0.140, 0.160,
#
     0.180,\ 0.200,\ 0.220,\ 0.240,\ 0.260,\ 0.280,\ 0.300,\ 0.320,\ 0.340,
#
     0.360, 0.380, 0.400, 0.420, 0.440, 0.460, 0.480, 0.500, 0.520,
#
     0.540, 0.560, 0.580, 0.600, 0.620, 0.640, 0.660, 0.680, 0.700,
#
     0.720, 0.740, 0.760, 0.780, 0.800, 0.820, 0.840, 0.860, 0.880,
#
     0.900, 0.920, 0.940, 0.960, 0.980, 1.000, 1.020, 1.040, 1.060,
     1.080, 1.100, 1.120, 1.140, 1.160, 1.180, 1.200, 1.220, 1.240,
#
     1.260, 1.280, 1.300, 1.320, 1.340, 1.360, 1.380, 1.400, 1.420,
#
     1.440, 1.460, 1.480, 1.500, 1.520, 1.540, 1.560, 1.580, 1.600,
#
     1.620, 1.640, 1.660, 1.680, 1.700, 1.720, 1.740, 1.760, 1.780/
    DATA (RIGPC(I), I=91,180)/
     1.800, 1.820, 1.840, 1.860, 1.880, 1.900, 1.920, 1.940, 1.960,
#
     1.980, 2.000, 2.020, 2.040, 2.060, 2.080, 2.100, 2.120, 2.140,
#
     2.160, 2.180, 2.200, 2.220, 2.240, 2.260, 2.280, 2.300, 2.320,
     2.340, 2.360, 2.380, 2.400, 2.420, 2.440, 2.460, 2.480, 2.500,
     2.520, 2.540, 2.560, 2.580, 2.600, 2.620, 2.640, 2.660, 2.680,
     2.700, 2.720, 2.740, 2.760, 2.780, 2.800, 2.820, 2.840, 2.860,
     2.880, 2.900, 2.920, 2.940, 2.960, 2.980, 3.000, 3.020, 3.040,
     3.060, 3.080, 3.100, 3.120, 3.140, 3.160, 3.180, 3.200, 3.220,
     3.240, 3.260, 3.280, 3.300, 3.320, 3.340, 3.360, 3.380, 3.400,
```

DATA (CUTOFF (32, J), J=1, 72)

```
3.420, 3.440, 3.460, 3.480, 3.500, 3.520, 3.540, 3.560, 3.580/
#
    DATA (RIGPC(I), I=181,270)/
     3.600, 3.620, 3.640, 3.660, 3.680, 3.700, 3.720, 3.740, 3.760,
#
     3.780, 3.800, 3.820, 3.840, 3.860, 3.880, 3.900, 3.920, 3.940,
     3.960, 3.980, 4.000, 4.020, 4.040, 4.060, 4.080, 4.100, 4.120,
#
     4.140, 4.160, 4.180, 4.200, 4.220, 4.240, 4.260, 4.280, 4.300,
#
     4.320, 4.340, 4.360, 4.380, 4.400, 4.420, 4.440, 4.460, 4.480,
#
     4.500, 4.520, 4.540, 4.560, 4.580, 4.600, 4.620, 4.640, 4.660,
#
     4.680, 4.700, 4.720, 4.740, 4.760, 4.780, 4.800, 4.820, 4.840,
#
     4.860, 4.880, 4.900, 4.920, 4.940, 4.960, 4.980, 5.000, 5.020,
#
     5.040, 5.060, 5.080, 5.100, 5.120, 5.140, 5.160, 5.180, 5.200,
#
     5.220, 5.240, 5.260, 5.280, 5.300, 5.320, 5.340, 5.360, 5.380/
#
    DATA (RIGPC(I), I=271, 360)/
     5.400, 5.420, 5.440, 5.460, 5.480, 5.500, 5.520, 5.540, 5.560,
     5.580, 5.600, 5.620, 5.640, 5.660, 5.680, 5.700, 5.720, 5.740,
#
     5.760, 5.780, 5.800, 5.820, 5.840, 5.860, 5.880, 5.900, 5.920,
#
     5.940, 5.960, 5.980, 6.000, 6.020, 6.040, 6.060, 6.080, 6.100,
#
     6.120, 6.140, 6.160, 6.180, 6.200, 6.220, 6.240, 6.260, 6.280,
#
     6.300, 6.320, 6.340, 6.360, 6.380, 6.400, 6.420, 6.440, 6.460,
#
     6.480, 6.500, 6.520, 6.540, 6.560, 6.580, 6.600, 6.620, 6.640,
#
     6.660, 6.680, 6.700, 6.720, 6.740, 6.760, 6.780, 6.800, 6.820,
#
     6.840, 6.860, 6.880, 6.900, 6.920, 6.940, 6.960, 6.980, 7.000,
#
     7.020, 7.040, 7.060, 7.080, 7.100, 7.120, 7.140, 7.160, 7.180/
    DATA (RIGPC(I), I=361,450)/
     7.200, 7.220, 7.240, 7.260, 7.280, 7.300, 7.320, 7.340, 7.360,
#
     7.380, 7.400, 7.420, 7.440, 7.460, 7.480, 7.500, 7.520, 7.540,
#
     7.560, 7.580, 7.600, 7.620, 7.640, 7.660, 7.680, 7.700, 7.720,
#
     7.740, 7.760, 7.780, 7.800, 7.820, 7.840, 7.860, 7.880, 7.900,
#
     7.920, 7.940, 7.960, 7.980, 8.000, 8.020, 8.040, 8.060, 8.080,
#
     8.100, 8.120, 8.140, 8.160, 8.180, 8.200, 8.220, 8.240, 8.260,
#
     8.280, 8.300, 8.320, 8.340, 8.360, 8.380, 8.400, 8.420, 8.440,
#
     8.460, 8.480, 8.500, 8.520, 8.540, 8.560, 8.580, 8.600, 8.620,
#
     8.640, 8.660, 8.680, 8.700, 8.720, 8.740, 8.760, 8.780, 8.800,
#
     8.820, 8.840, 8.860, 8.880, 8.900, 8.920, 8.940, 8.960, 8.980/
#
    DATA (RIGPC(I), I=451,540)/
     9.000, 9.020, 9.040, 9.060, 9.080, 9.100, 9.120, 9.140, 9.160,
#
     9.180, 9.200, 9.220, 9.240, 9.260, 9.280, 9.300, 9.320, 9.340,
#
     9.360, 9.380, 9.400, 9.420, 9.440, 9.460, 9.480, 9.500, 9.520,
#
     9.540, 9.560, 9.580, 9.600, 9.620, 9.640, 9.660, 9.680, 9.700,
#
     9.720, 9.740, 9.760, 9.780, 9.800, 9.820, 9.840, 9.860, 9.880,
#
     9.900, 9.920, 9.940, 9.960, 9.980,10.000,10.020,10.040,10.060,
#
     10.080, 10.100, 10.120, 10.140, 10.160, 10.180, 10.200, 10.220, 10.240,
#
     10.260,10.280,10.300,10.320,10.340,10.360,10.380,10.400,10.420,
#
     10.440,10.460,10.480,10.500,10.520,10.540,10.560,10.580,10.600,
#
     10.620,10.640,10.660,10.680,10.700,10.720,10.740,10.760,10.780/
#
     DATA (RIGPC(I), I=541,630)/
     10.800,10.820,10.840,10.860,10.880,10.900,10.920,10.940,10.960,
#
     10.980,11.000,11.020,11.040,11.060,11.080,11.100,11.120,11.140,
 #
     11.160,11.180,11.200,11.220,11.240,11.260,11.280,11.300,11.320,
 #
     11.340,11.360,11.380,11.400,11.420,11.440,11.460,11.480,11.500,
 #
     11.520,11.540,11.560,11.580,11.600,11.620,11.640,11.660,11.680,
 #
     11.700,11.720,11.740,11.760,11.780,11.800,11.820,11.840,11.860,
     11.880,11.900,11.920,11.940,11.960,11.980,12.000,12.020,12.040,
 #
     12.060, 12.080, 12.100, 12.120, 12.140, 12.160, 12.180, 12.200, 12.220,
     12.240, 12.260, 12.280, 12.300, 12.320, 12.340, 12.360, 12.380, 12.400,
     12.420,12.440,12.460,12.480,12.500,12.520,12.540,12.560,12.580/
     DATA (RIGPC(I), I=631,720)/
     12.600,12.620,12.640,12.660,12.680,12.700,12.720,12.740,12.760,
     12.780,12.800,12.820,12.840,12.860,12.880,12.900,12.920,12.940,
     12.960,12.980,13.000,13.020,13.040,13.060,13.080,13.100,13.120,
```

```
13.140,13.160,13.180,13.200,13.220,13.240,13.260,13.280,13.300,
    13.320,13.340,13.360,13.380,13.400,13.420,13.440,13.460,13.480,
#
#
    13.500, 13.520, 13.540, 13.560, 13.580, 13.600, 13.620, 13.640, 13.660,
#
    13.680, 13.700, 13.720, 13.740, 13.760, 13.780, 13.800, 13.820, 13.840,
    13.860,13.880,13.900,13.920,13.940,13.960,13.980,14.000,14.020,
    14.040,14.060,14.080,14.100,14.120,14.140,14.160,14.180,14.200,
    14.220,14.240,14.260,14.280,14.300,14.320,14.340,14.360,14.380/
    DATA (RIGPC(I), I=721,810)/
    14.400,14.420,14.440,14.460,14.480,14.500,14.520,14.540,14.560,
#
#
    14.580, 14.600, 14.620, 14.640, 14.660, 14.680, 14.700, 14.720, 14.740,
#
    14.760, 14.780, 14.800, 14.820, 14.840, 14.860, 14.880, 14.900, 14.920,
    14.940,14.960,14.980,15.000,15.020,15.040,15.060,15.080,15.100,
#
    15.120,15.140,15.160,15.180,15.200,15.220,15.240,15.260,15.280,
    15.300,15.320,15.340,15.360,15.380,15.400,15.420,15.440,15.460,
    15.480, 15.500, 15.520, 15.540, 15.560, 15.580, 15.600, 15.620, 15.640,
    15.660, 15.680, 15.700, 15.720, 15.740, 15.760, 15.780, 15.800, 15.820,
#
    15.840,15.860,15.880,15.900,15.920,15.940,15.960,15.980,16.000,
    16.020, 16.040, 16.060, 16.080, 16.100, 16.120, 16.140, 16.160, 16.180/
    DATA (RIGPC(I), I=811,900)/
#
    16.200,16.220,16.240,16.260,16.280,16.300,16.320,16.340,16.360,
    16.380,16.400,16.420,16.440,16.460,16.480,16.500,16.520,16.540,
#
    16.560,16.580,16.600,16.620,16.640,16.660,16.680,16.700,16.720,
    16.740,16.760,16.780,16.800,16.820,16.840,16.860,16.880,16.900,
#
    16.920, 16.940, 16.960, 16.980, 17.000, 17.020, 17.040, 17.060, 17.080,
#
    17.100,17.120,17.140,17.160,17.180,17.200,17.220,17.240,17.260,
#
    17.280,17.300,17.320,17.340,17.360,17.380,17.400,17.420,17.440,
#
    17.460,17.480,17.500,17.520,17.540,17.560,17.580,17.600,17.620,
    17.640,17.660,17.680,17.700,17.720,17.740,17.760,17.780,17.800,
#
#
    17.820,17.840,17.860,17.880,17.900,17.920,17.940,17.960,17.980/
    DATA (RIGPC(I), I=901, 990) /
#
    18.000, 18.020, 18.040, 18.060, 18.080, 18.100, 18.120, 18.140, 18.160,
    18.180, 18.200, 18.220, 18.240, 18.260, 18.280, 18.300, 18.320, 18.340,
    18.360,18.380,18.400,18.420,18.440,18.460,18.480,18.500,18.520,
#
#
    18.540, 18.560, 18.580, 18.600, 18.620, 18.640, 18.660, 18.680, 18.700,
#
    18.720,18.740,18.760,18.780,18.800,18.820,18.840,18.860,18.880,
    18.900,18.920,18.940,18.960,18.980,19.000,19.020,19.040,19.060,
#
    19.080,19.100,19.120,19.140,19.160,19.180,19.200,19.220,19.240,
    19.260, 19.280, 19.300, 19.320, 19.340, 19.360, 19.380, 19.400, 19.420,
    19.440,19.460,19.480,19.500,19.520,19.540,19.560,19.580,19.600,
    19.620,19.640,19.660,19.680,19.700,19.720,19.740,19.760,19.780/
    DATA (RIGPC(I), I=991, Nrigs)/
#
    19.800,19.820,19.840,19.860,19.880,19.900,19.920,19.940,19.960,
#
    19.980,20.000/
    DATA (PCGTF1(I), I=1,50)/
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.2139E-03,0.5824E-03,0.1253E-02,0.2217E-02,
#
    0.3307E-02,0.4387E-02,0.5475E-02,0.6627E-02,0.7837E-02,
    0.9039E-02,0.1019E-01,0.1139E-01,0.1273E-01,0.1424E-01,
    0.1583E-01,0.1741E-01,0.1889E-01,0.2021E-01,0.2132E-01,
#
    0.2230E-01,0.2319E-01,0.2404E-01,0.2491E-01,0.2583E-01,
#
    0.2678E-01,0.2778E-01,0.2881E-01,0.2987E-01,0.3096E-01,
    0.3209E-01,0.3323E-01,0.3440E-01,0.3558E-01,0.3678E-01,
    0.3799E-01,0.3921E-01,0.4044E-01,0.4167E-01,0.4290E-01,
    0.4413E-01,0.4535E-01,0.4657E-01,0.4777E-01,0.4896E-01/
   DATA (PCGTF1(I), I=51,100)/
    0.5014E-01,0.5129E-01,0.5242E-01,0.5352E-01,0.5460E-01,
    0.5565E-01,0.5668E-01,0.5769E-01,0.5867E-01,0.5963E-01,
    0.6057E-01,0.6149E-01,0.6239E-01,0.6327E-01,0.6414E-01,
    0.6500E-01,0.6584E-01,0.6666E-01,0.6748E-01,0.6828E-01,
```

```
0.6908E-01,0.6986E-01,0.7064E-01,0.7141E-01,0.7218E-01,
#
    0.7294E-01,0.7370E-01,0.7446E-01,0.7522E-01,0.7597E-01,
#
    0.7673E-01,0.7749E-01,0.7825E-01,0.7902E-01,0.7979E-01,
#
    0.8057E-01,0.8135E-01,0.8215E-01,0.8295E-01,0.8376E-01,
#
    0.8459E-01,0.8543E-01,0.8628E-01,0.8715E-01,0.8803E-01,
#
    0.8894E-01,0.8985E-01,0.9079E-01,0.9175E-01,0.9273E-01/
#
    DATA (PCGTF1(I), I=101, 150)/
    0.9373E-01,0.9475E-01,0.9579E-01,0.9684E-01,0.9791E-01,
#
    0.9900E-01,0.1001E+00,0.1012E+00,0.1024E+00,0.1035E+00,
#
    0.1047E+00,0.1059E+00,0.1071E+00,0.1083E+00,0.1095E+00,
#
    0.1107E+00,0.1119E+00,0.1132E+00,0.1144E+00,0.1157E+00,
#
    0.1170E+00,0.1183E+00,0.1195E+00,0.1208E+00,0.1221E+00,
#
    0.1234E+00,0.1247E+00,0.1261E+00,0.1274E+00,0.1287E+00,
#
    0.1300E+00,0.1313E+00,0.1326E+00,0.1340E+00,0.1353E+00,
#
    0.1366E+00,0.1379E+00,0.1392E+00,0.1406E+00,0.1419E+00,
#
    0.1432E+00,0.1445E+00,0.1458E+00,0.1471E+00,0.1483E+00,
#
    0.1496E+00,0.1509E+00,0.1522E+00,0.1534E+00,0.1546E+00/
    DATA (PCGTF1(I), I=151,200)/
    0.1559E+00,0.1571E+00,0.1583E+00,0.1595E+00,0.1607E+00,
#
    0.1619E+00,0.1631E+00,0.1643E+00,0.1654E+00,0.1666E+00,
#
    0.1677E+00,0.1689E+00,0.1700E+00,0.1711E+00,0.1723E+00,
#
    0.1734E+00,0.1745E+00,0.1756E+00,0.1766E+00,0.1777E+00,
#
    0.1788E+00,0.1799E+00,0.1809E+00,0.1820E+00,0.1830E+00,
#
    0.1840E+00,0.1851E+00,0.1861E+00,0.1871E+00,0.1881E+00,
#
    0.1891E+00,0.1901E+00,0.1911E+00,0.1921E+00,0.1930E+00,
#
    0.1940E+00,0.1950E+00,0.1959E+00,0.1969E+00,0.1978E+00,
#
    0.1987E+00,0.1997E+00,0.2006E+00,0.2015E+00,0.2024E+00,
#
    0.2033E+00,0.2042E+00,0.2051E+00,0.2060E+00,0.2069E+00/
#
    DATA (PCGTF1(I), I=201, 250)/
    0.2078E+00,0.2087E+00,0.2095E+00,0.2104E+00,0.2113E+00,
#
    0.2121E+00,0.2130E+00,0.2138E+00,0.2147E+00,0.2155E+00,
#
    0.2163E+00,0.2172E+00,0.2180E+00,0.2188E+00,0.2196E+00,
#
    0.2204E+00,0.2212E+00,0.2220E+00,0.2228E+00,0.2236E+00,
#
    0.2244E+00,0.2252E+00,0.2260E+00,0.2268E+00,0.2276E+00,
#
    0.2283E+00,0.2291E+00,0.2299E+00,0.2306E+00,0.2314E+00,
#
    0.2322E+00,0.2329E+00,0.2337E+00,0.2344E+00,0.2352E+00,
#
    0.2359E+00,0.2367E+00,0.2374E+00,0.2381E+00,0.2389E+00,
#
    0.2396E+00,0.2403E+00,0.2411E+00,0.2418E+00,0.2425E+00,
#
    0.2432E+00,0.2440E+00,0.2447E+00,0.2454E+00,0.2461E+00/
#
    DATA (PCGTF1(I), I=251,300)/
    0.2468E+00,0.2475E+00,0.2483E+00,0.2490E+00,0.2497E+00,
#
    0.2504E+00,0.2511E+00,0.2518E+00,0.2525E+00,0.2532E+00,
#
    0.2539E+00,0.2546E+00,0.2553E+00,0.2560E+00,0.2567E+00,
#
    0.2574E+00,0.2581E+00,0.2588E+00,0.2595E+00,0.2602E+00,
#
    0.2609E+00,0.2616E+00,0.2622E+00,0.2629E+00,0.2636E+00,
#
    0.2643E+00,0.2650E+00,0.2657E+00,0.2664E+00,0.2670E+00,
#
    0.2677E+00,0.2684E+00,0.2691E+00,0.2698E+00,0.2704E+00,
#
    0.2711E+00,0.2718E+00,0.2725E+00,0.2731E+00,0.2738E+00,
#
    0.2745E+00,0.2752E+00,0.2758E+00,0.2765E+00,0.2772E+00,
#
#
    0.2779E+00,0.2785E+00,0.2792E+00,0.2799E+00,0.2805E+00/
    DATA (PCGTF1(I), I=301,350)/
    0.2812E+00,0.2819E+00,0.2826E+00,0.2832E+00,0.2839E+00,
#
    0.2846E+00,0.2852E+00,0.2859E+00,0.2866E+00,0.2872E+00,
#
    0.2879E+00,0.2886E+00,0.2892E+00,0.2899E+00,0.2906E+00,
#
    0.2912E+00,0.2919E+00,0.2926E+00,0.2932E+00,0.2939E+00,
#
#
    0.2946E+00,0.2952E+00,0.2959E+00,0.2966E+00,0.2972E+00,
    0.2979E+00,0.2985E+00,0.2992E+00,0.2999E+00,0.3005E+00,
#
    0.3012E+00,0.3019E+00,0.3025E+00,0.3032E+00,0.3039E+00,
#
    0.3046E+00,0.3052E+00,0.3059E+00,0.3066E+00,0.3072E+00,
#
    0.3079E+00,0.3086E+00,0.3092E+00,0.3099E+00,0.3106E+00,
```

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0.3113E+00,0.3119E+00,0.3126E+00,0.3133E+00,0.3139E+00/
    DATA (PCGTF1(I), I=351, 400)/
    0.3146E+00,0.3153E+00,0.3160E+00,0.3167E+00,0.3173E+00,
#
    0.3180E+00,0.3187E+00,0.3194E+00,0.3200E+00,0.3207E+00,
    0.3214E+00,0.3221E+00,0.3228E+00,0.3235E+00,0.3241E+00,
    0.3248E+00,0.3255E+00,0.3262E+00,0.3269E+00,0.3276E+00,
#
    0.3283E+00,0.3290E+00,0.3297E+00,0.3304E+00,0.3311E+00,
#
    0.3318E+00,0.3324E+00,0.3331E+00,0.3338E+00,0.3345E+00,
    0.3352E+00,0.3360E+00,0.3367E+00,0.3374E+00,0.3381E+00,
#
    0.3388E+00,0.3395E+00,0.3402E+00,0.3409E+00,0.3416E+00,
#
    0.3423E+00,0.3430E+00,0.3437E+00,0.3445E+00,0.3452E+00,
#
    0.3459E+00,0.3466E+00,0.3473E+00,0.3480E+00,0.3488E+00/
    DATA (PCGTF1(I), I=401,450)/
    0.3495E+00,0.3502E+00,0.3509E+00,0.3516E+00,0.3524E+00,
#
    0.3531E+00,0.3538E+00,0.3545E+00,0.3552E+00,0.3560E+00,
#
    0.3567E+00,0.3574E+00,0.3581E+00,0.3589E+00,0.3596E+00,
#
    0.3603E+00,0.3610E+00,0.3618E+00,0.3625E+00,0.3632E+00,
#
    0.3639E+00,0.3647E+00,0.3654E+00,0.3661E+00,0.3668E+00,
#
    0.3676E+00,0.3683E+00,0.3690E+00,0.3697E+00,0.3705E+00,
#
    0.3712E+00,0.3719E+00,0.3726E+00,0.3734E+00,0.3741E+00,
#
    0.3748E+00,0.3755E+00,0.3763E+00,0.3770E+00,0.3777E+00,
#
    0.3784E+00,0.3792E+00,0.3799E+00,0.3806E+00,0.3813E+00,
#
    0.3821E+00,0.3828E+00,0.3835E+00,0.3842E+00,0.3850E+00/
#
    DATA (PCGTF1(I), I=451,500)/
    0.3857E+00,0.3864E+00,0.3871E+00,0.3878E+00,0.3886E+00,
#
    0.3893E+00,0.3900E+00,0.3907E+00,0.3914E+00,0.3921E+00,
#
    0.3929E+00,0.3936E+00,0.3943E+00,0.3950E+00,0.3957E+00,
#
    0.3964E+00,0.3971E+00,0.3978E+00,0.3985E+00,0.3993E+00,
#
    0.4000E+00,0.4007E+00,0.4014E+00,0.4021E+00,0.4028E+00,
#
    0.4035E+00,0.4042E+00,0.4049E+00,0.4056E+00,0.4063E+00,
#
    0.4070E+00,0.4077E+00,0.4084E+00,0.4091E+00,0.4097E+00,
#
    0.4104E+00,0.4111E+00,0.4118E+00,0.4125E+00,0.4132E+00,
#
    0.4139E+00,0.4146E+00,0.4152E+00,0.4159E+00,0.4166E+00,
#
    0.4173E+00,0.4179E+00,0.4186E+00,0.4193E+00,0.4200E+00/
#
    DATA (PCGTF1(I), I=501,550)/
    0.4206E+00,0.4209E+00,0.4212E+00,0.4215E+00,0.4218E+00,
#
    0.4220E+00,0.4223E+00,0.4226E+00,0.4229E+00,0.4232E+00,
#
    0.4234E+00,0.4237E+00,0.4240E+00,0.4243E+00,0.4246E+00,
    0.4249E+00,0.4251E+00,0.4254E+00,0.4257E+00,0.4260E+00,
#
    0.4263E+00,0.4266E+00,0.4268E+00,0.4271E+00,0.4274E+00,
#
    0.4277E+00,0.4280E+00,0.4283E+00,0.4286E+00,0.4288E+00,
#
    0.4291E+00,0.4294E+00,0.4297E+00,0.4300E+00,0.4303E+00,
    0.4306E+00,0.4309E+00,0.4311E+00,0.4314E+00,0.4317E+00,
#
    0.4320E+00,0.4323E+00,0.4326E+00,0.4329E+00,0.4332E+00,
    0.4334E+00,0.4337E+00,0.4340E+00,0.4343E+00,0.4346E+00/
    DATA (PCGTF1(I), I=551,600)/
    0.4349E+00,0.4352E+00,0.4355E+00,0.4358E+00,0.4361E+00,
#
    0.4363E+00,0.4366E+00,0.4369E+00,0.4372E+00,0.4375E+00,
#
    0.4378E+00,0.4381E+00,0.4384E+00,0.4387E+00,0.4390E+00,
#
    0.4393E+00,0.4396E+00,0.4399E+00,0.4401E+00,0.4404E+00,
#
    0.4407E+00,0.4410E+00,0.4413E+00,0.4416E+00,0.4419E+00,
#
    0.4422E+00,0.4425E+00,0.4428E+00,0.4431E+00,0.4434E+00,
#
    0.4437E+00,0.4440E+00,0.4443E+00,0.4446E+00,0.4449E+00,
    0.4452E+00,0.4455E+00,0.4458E+00,0.4461E+00,0.4464E+00,
#
    0.4467E+00,0.4469E+00,0.4472E+00,0.4475E+00,0.4478E+00,
    0.4481E+00,0.4484E+00,0.4487E+00,0.4490E+00,0.4493E+00/
#
    DATA (PCGTF1(I), I=601,650)/
    0.4496E+00,0.4499E+00,0.4502E+00,0.4505E+00,0.4508E+00,
#
    0.4511E+00,0.4514E+00,0.4517E+00,0.4520E+00,0.4523E+00,
     0.4526E+00,0.4530E+00,0.4533E+00,0.4536E+00,0.4539E+00,
```

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0.4542E+00,0.4545E+00,0.4548E+00,0.4551E+00,0.4554E+00,
    0.4557E+00,0.4560E+00,0.4563E+00,0.4566E+00,0.4569E+00,
#
    0.4572E+00,0.4575E+00,0.4578E+00,0.4581E+00,0.4584E+00,
#
    0.4587E+00,0.4590E+00,0.4593E+00,0.4596E+00,0.4600E+00,
#
    0.4603E+00,0.4606E+00,0.4609E+00,0.4612E+00,0.4615E+00,
    0.4618E+00,0.4621E+00,0.4624E+00,0.4627E+00,0.4630E+00,
#
    0.4633E+00,0.4636E+00,0.4640E+00,0.4643E+00,0.4646E+00/
#
    DATA (PCGTF1(I), I=651,700)/
    0.4649E+00,0.4652E+00,0.4655E+00,0.4658E+00,0.4661E+00,
#
    0.4664E+00,0.4668E+00,0.4671E+00,0.4674E+00,0.4677E+00,
#
    0.4680E+00,0.4683E+00,0.4686E+00,0.4689E+00,0.4692E+00,
#
    0.4696E+00,0.4699E+00,0.4702E+00,0.4705E+00,0.4708E+00,
#
    0.4711E+00,0.4714E+00,0.4718E+00,0.4721E+00,0.4724E+00,
#
    0.4727E+00,0.4730E+00,0.4733E+00,0.4736E+00,0.4740E+00,
#
    0.4743E+00,0.4746E+00,0.4749E+00,0.4752E+00,0.4755E+00,
    0.4759E+00,0.4762E+00,0.4765E+00,0.4768E+00,0.4771E+00,
#
    0.4775E+00,0.4778E+00,0.4781E+00,0.4784E+00,0.4787E+00,
#
    0.4791E+00,0.4794E+00,0.4797E+00,0.4800E+00,0.4803E+00/
#
    DATA (PCGTF1(I), I=701,750)/
    0.4807E+00,0.4810E+00,0.4813E+00,0.4816E+00,0.4819E+00,
#
    0.4823E+00,0.4826E+00,0.4829E+00,0.4832E+00,0.4835E+00,
#
    0.4839E+00,0.4842E+00,0.4845E+00,0.4848E+00,0.4852E+00,
#
    0.4855E+00,0.4858E+00,0.4861E+00,0.4865E+00,0.4868E+00,
#
    0.4871E+00,0.4874E+00,0.4878E+00,0.4881E+00,0.4884E+00,
#
    0.4887E+00,0.4891E+00,0.4894E+00,0.4897E+00,0.4900E+00,
#
    0.4904E+00,0.4907E+00,0.4910E+00,0.4913E+00,0.4917E+00,
#
    0.4920E+00,0.4923E+00,0.4927E+00,0.4930E+00,0.4933E+00,
#
    0.4936E+00,0.4940E+00,0.4943E+00,0.4946E+00,0.4950E+00,
#
    0.4953E+00,0.4956E+00,0.4960E+00,0.4963E+00,0.4966E+00/
#
    DATA (PCGTF1(I), I=751,800)/
    0.4969E+00,0.4973E+00,0.4976E+00,0.4979E+00,0.4983E+00,
#
    0.4986E+00,0.4989E+00,0.4993E+00,0.4996E+00,0.4999E+00,
#
    0.5003E+00,0.5006E+00,0.5009E+00,0.5013E+00,0.5016E+00,
#
    0.5019E+00,0.5023E+00,0.5026E+00,0.5030E+00,0.5033E+00,
#
    0.5036E+00,0.5040E+00,0.5043E+00,0.5046E+00,0.5050E+00,
#
    0.5053E+00,0.5056E+00,0.5060E+00,0.5063E+00,0.5067E+00,
#
    0.5070E+00,0.5073E+00,0.5077E+00,0.5080E+00,0.5083E+00,
#
    0.5087E+00,0.5090E+00,0.5094E+00,0.5097E+00,0.5100E+00,
    0.5104E+00,0.5107E+00,0.5111E+00,0.5114E+00,0.5117E+00,
#
    0.5121E+00,0.5124E+00,0.5128E+00,0.5131E+00,0.5135E+00/
#
    DATA (PCGTF1(I), I=801,850)/
    0.5138E+00,0.5141E+00,0.5145E+00,0.5148E+00,0.5152E+00,
#
    0.5155E+00,0.5159E+00,0.5162E+00,0.5165E+00,0.5169E+00,
#
    0.5172E+00,0.5176E+00,0.5179E+00,0.5183E+00,0.5186E+00,
#
    0.5190E+00,0.5193E+00,0.5197E+00,0.5200E+00,0.5204E+00,
#
    0.5207E+00,0.5210E+00,0.5214E+00,0.5217E+00,0.5221E+00,
    0.5224E+00,0.5228E+00,0.5231E+00,0.5235E+00,0.5238E+00,
#
    0.5242E+00,0.5245E+00,0.5249E+00,0.5252E+00,0.5256E+00,
#
    0.5259E+00,0.5263E+00,0.5266E+00,0.5270E+00,0.5273E+00,
#
    0.5277E+00,0.5280E+00,0.5284E+00,0.5287E+00,0.5291E+00,
    0.5295E+00,0.5298E+00,0.5302E+00,0.5305E+00,0.5309E+00/
#
    DATA (PCGTF1(I), I=851,900)/
    0.5312E+00,0.5316E+00,0.5319E+00,0.5323E+00,0.5326E+00,
#
    0.5330E+00,0.5334E+00,0.5337E+00,0.5341E+00,0.5344E+00,
    0.5348E+00,0.5351E+00,0.5355E+00,0.5358E+00,0.5362E+00,
#
    0.5366E+00,0.5369E+00,0.5373E+00,0.5376E+00,0.5380E+00,
#
    0.5384E+00,0.5387E+00,0.5391E+00,0.5394E+00,0.5398E+00,
#
    0.5402E+00,0.5405E+00,0.5409E+00,0.5412E+00,0.5416E+00,
    0.5420E+00,0.5423E+00,0.5427E+00,0.5430E+00,0.5434E+00,
     0.5438E+00,0.5441E+00,0.5445E+00,0.5449E+00,0.5452E+00,
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0.5456E+00,0.5459E+00,0.5463E+00,0.5467E+00,0.5470E+00,
    0.5474E+00,0.5478E+00,0.5481E+00,0.5485E+00,0.5489E+00/
#
    DATA (PCGTF1(I), I=901,950)/
    0.5492E+00,0.5496E+00,0.5500E+00,0.5503E+00,0.5507E+00,
    0.5511E+00,0.5514E+00,0.5518E+00,0.5522E+00,0.5525E+00,
    0.5529E+00,0.5533E+00,0.5536E+00,0.5540E+00,0.5544E+00,
#
#
    0.5548E+00,0.5551E+00,0.5555E+00,0.5559E+00,0.5562E+00,
    0.5566E+00,0.5570E+00,0.5574E+00,0.5577E+00,0.5581E+00,
    0.5585E+00,0.5588E+00,0.5592E+00,0.5596E+00,0.5600E+00,
#
#
    0.5603E+00,0.5607E+00,0.5611E+00,0.5615E+00,0.5618E+00,
    0.5622E+00,0.5626E+00,0.5630E+00,0.5633E+00,0.5637E+00,
#
    0.5641E+00,0.5645E+00,0.5648E+00,0.5652E+00,0.5656E+00,
#
    0.5660E+00,0.5663E+00,0.5667E+00,0.5671E+00,0.5675E+00/
    DATA (PCGTF1(I), I=951, Nrigs)/
    0.5679E+00,0.5682E+00,0.5686E+00,0.5690E+00,0.5694E+00,
#
    0.5698E+00,0.5701E+00,0.5705E+00,0.5709E+00,0.5713E+00,
#
    0.5717E+00,0.5720E+00,0.5724E+00,0.5728E+00,0.5732E+00,
#
#
    0.5736E+00,0.5740E+00,0.5743E+00,0.5747E+00,0.5751E+00,
    0.5755E+00,0.5759E+00,0.5763E+00,0.5766E+00,0.5770E+00,
#
    0.5774E+00,0.5778E+00,0.5782E+00,0.5786E+00,0.5789E+00,
#
    0.5793E+00,0.5797E+00,0.5801E+00,0.5805E+00,0.5809E+00,
#
    0.5813E+00,0.5817E+00,0.5820E+00,0.5824E+00,0.5828E+00,
    0.5832E+00,0.5836E+00,0.5840E+00,0.5844E+00,0.5848E+00,
    0.5852E+00,0.5856E+00,0.5859E+00,0.5863E+00,0.5867E+00,
#
#
    0.5871E+00/
   DATA (PCGTF2(I), I=1,50)/
    0.4848E-01,0.4962E-01,0.5075E-01,0.5189E-01,0.5303E-01,
#
#
    0.5417E-01,0.5533E-01,0.5646E-01,0.5754E-01,0.5861E-01,
    0.5976E-01,0.6109E-01,0.6272E-01,0.6472E-01,0.6689E-01,
#
    0.6874E-01,0.6991E-01,0.7067E-01,0.7145E-01,0.7242E-01,
    0.7351E-01,0.7463E-01,0.7570E-01,0.7664E-01,0.7742E-01,
#
#
    0.7809E-01,0.7869E-01,0.7927E-01,0.7987E-01,0.8050E-01,
#
    0.8118E-01,0.8190E-01,0.8265E-01,0.8343E-01,0.8425E-01,
    0.8510E-01,0.8597E-01,0.8687E-01,0.8779E-01,0.8873E-01,
#
    0.8970E-01,0.9068E-01,0.9168E-01,0.9269E-01,0.9371E-01,
    0.9474E-01,0.9578E-01,0.9682E-01,0.9787E-01,0.9892E-01/
#
   DATA (PCGTF2(I), I=51, 100)/
#
    0.9997E-01,0.1010E+00,0.1021E+00,0.1031E+00,0.1041E+00,
#
    0.1051E+00,0.1062E+00,0.1072E+00,0.1082E+00,0.1092E+00,
#
    0.1102E+00,0.1111E+00,0.1121E+00,0.1131E+00,0.1141E+00,
#
    0.1151E+00,0.1160E+00,0.1170E+00,0.1179E+00,0.1189E+00,
#
    0.1199E+00,0.1208E+00,0.1218E+00,0.1227E+00,0.1237E+00,
#
    0.1246E+00,0.1256E+00,0.1266E+00,0.1275E+00,0.1285E+00,
#
    0.1294E+00,0.1304E+00,0.1313E+00,0.1323E+00,0.1333E+00,
#
    0.1343E+00,0.1352E+00,0.1362E+00,0.1372E+00,0.1382E+00,
    0.1392E+00,0.1402E+00,0.1412E+00,0.1422E+00,0.1432E+00,
#
    0.1442E+00,0.1453E+00,0.1463E+00,0.1474E+00,0.1484E+00/
   DATA (PCGTF2(I), I=101, 150)/
#
    0.1495E+00,0.1505E+00,0.1516E+00,0.1527E+00,0.1538E+00,
#
    0.1549E+00,0.1560E+00,0.1571E+00,0.1582E+00,0.1593E+00,
#
    0.1604E+00,0.1616E+00,0.1627E+00,0.1638E+00,0.1650E+00,
#
    0.1661E+00,0.1672E+00,0.1684E+00,0.1695E+00,0.1707E+00,
#
    0.1718E+00,0.1729E+00,0.1741E+00,0.1752E+00,0.1764E+00,
    0.1775E+00,0.1787E+00,0.1798E+00,0.1810E+00,0.1821E+00,
    0.1832E+00,0.1844E+00,0.1855E+00,0.1866E+00,0.1878E+00,
#
#
    0.1889E+00,0.1900E+00,0.1911E+00,0.1922E+00,0.1933E+00,
#
    0.1944E+00,0.1955E+00,0.1966E+00,0.1977E+00,0.1988E+00,
    0.1998E+00,0.2009E+00,0.2020E+00,0.2030E+00,0.2040E+00/
   DATA (PCGTF2(I), I=151,200)/
    0.2051E+00,0.2061E+00,0.2071E+00,0.2081E+00,0.2091E+00,
```

```
0.2101E+00,0.2111E+00,0.2121E+00,0.2130E+00,0.2140E+00,
    0.2149E+00,0.2159E+00,0.2168E+00,0.2178E+00,0.2187E+00,
    0.2196E+00,0.2205E+00,0.2214E+00,0.2223E+00,0.2232E+00,
    0.2241E+00,0.2250E+00,0.2258E+00,0.2267E+00,0.2276E+00,
    0.2284E+00,0.2293E+00,0.2301E+00,0.2310E+00,0.2318E+00,
#
    0.2326E+00,0.2334E+00,0.2343E+00,0.2351E+00,0.2359E+00,
#
    0.2367E+00,0.2375E+00,0.2382E+00,0.2390E+00,0.2398E+00,
    0.2406E+00,0.2414E+00,0.2421E+00,0.2429E+00,0.2436E+00,
    0.2444E+00,0.2451E+00,0.2459E+00,0.2466E+00,0.2473E+00/
    DATA (PCGTF2(I), I=201, 250)/
    0.2481E+00,0.2488E+00,0.2495E+00,0.2502E+00,0.2509E+00,
#
    0.2517E+00,0.2524E+00,0.2531E+00,0.2538E+00,0.2545E+00,
    0.2551E+00,0.2558E+00,0.2565E+00,0.2572E+00,0.2579E+00,
    0.2586E+00,0.2592E+00,0.2599E+00,0.2606E+00,0.2612E+00,
    0.2619E+00,0.2626E+00,0.2632E+00,0.2639E+00,0.2645E+00,
#
    0.2652E+00,0.2658E+00,0.2665E+00,0.2671E+00,0.2678E+00,
#
    0.2684E+00,0.2690E+00,0.2697E+00,0.2703E+00,0.2709E+00,
    0.2716E+00,0.2722E+00,0.2728E+00,0.2735E+00,0.2741E+00,
    0.2747E+00,0.2753E+00,0.2760E+00,0.2766E+00,0.2772E+00,
#
    0.2778E+00,0.2785E+00,0.2791E+00,0.2797E+00,0.2803E+00/
    DATA (PCGTF2(I), I=251,300)/
#
    0.2810E+00,0.2816E+00,0.2822E+00,0.2828E+00,0.2834E+00,
    0.2840E+00,0.2847E+00,0.2853E+00,0.2859E+00,0.2865E+00,
#
#
    0.2871E+00,0.2878E+00,0.2884E+00,0.2890E+00,0.2896E+00,
#
    0.2902E+00,0.2908E+00,0.2915E+00,0.2921E+00,0.2927E+00,
    0.2933E+00,0.2939E+00,0.2945E+00,0.2951E+00,0.2957E+00,
    0.2964E+00,0.2970E+00,0.2976E+00,0.2982E+00,0.2988E+00,
#
    0.2994E+00,0.3000E+00,0.3006E+00,0.3012E+00,0.3018E+00,
    0.3024E+00,0.3030E+00,0.3036E+00,0.3042E+00,0.3048E+00,
#
    0.3054E+00,0.3060E+00,0.3066E+00,0.3072E+00,0.3078E+00,
    0.3084E+00,0.3090E+00,0.3096E+00,0.3102E+00,0.3108E+00/
    DATA (PCGTF2(I), I=301, 350)/
    0.3114E+00,0.3120E+00,0.3126E+00,0.3132E+00,0.3138E+00,
#
    0.3144E+00,0.3150E+00,0.3155E+00,0.3161E+00,0.3167E+00,
    0.3173E+00,0.3179E+00,0.3185E+00,0.3190E+00,0.3196E+00,
    0.3202E+00,0.3208E+00,0.3213E+00,0.3219E+00,0.3225E+00,
#
    0.3231E+00,0.3236E+00,0.3242E+00,0.3248E+00,0.3253E+00,
#
    0.3259E+00,0.3265E+00,0.3270E+00,0.3276E+00,0.3282E+00,
    0.3287E+00,0.3293E+00,0.3298E+00,0.3304E+00,0.3310E+00,
    0.3315E+00,0.3321E+00,0.3326E+00,0.3332E+00,0.3337E+00,
    0.3343E+00,0.3348E+00,0.3353E+00,0.3359E+00,0.3364E+00,
#
    0.3370E+00,0.3375E+00,0.3380E+00,0.3386E+00,0.3391E+00/
   DATA (PCGTF2(I), I=351,400)/
    0.3396E+00,0.3402E+00,0.3407E+00,0.3412E+00,0.3417E+00,
    0.3423E+00,0.3428E+00,0.3433E+00,0.3438E+00,0.3444E+00,
    0.3449E+00,0.3454E+00,0.3459E+00,0.3464E+00,0.3469E+00,
#
    0.3474E+00,0.3479E+00,0.3484E+00,0.3489E+00,0.3494E+00,
    0.3499E+00,0.3504E+00,0.3509E+00,0.3514E+00,0.3519E+00,
    0.3524E+00,0.3529E+00,0.3534E+00,0.3538E+00,0.3543E+00,
    0.3548E+00,0.3553E+00,0.3557E+00,0.3562E+00,0.3567E+00,
    0.3572E+00,0.3576E+00,0.3581E+00,0.3586E+00,0.3590E+00,
#
   0.3595E+00,0.3599E+00,0.3604E+00,0.3609E+00,0.3613E+00,
    0.3618E+00,0.3622E+00,0.3627E+00,0.3631E+00,0.3636E+00/
   DATA (PCGTF2(I), I=401, 450)/
   0.3640E+00,0.3645E+00,0.3649E+00,0.3654E+00,0.3658E+00,
   0.3662E+00,0.3667E+00,0.3671E+00,0.3676E+00,0.3680E+00,
   0.3684E+00,0.3689E+00,0.3693E+00,0.3697E+00,0.3702E+00,
    0.3706E+00,0.3710E+00,0.3714E+00,0.3719E+00,0.3723E+00,
   0.3727E+00,0.3731E+00,0.3736E+00,0.3740E+00,0.3744E+00,
    0.3748E+00,0.3752E+00,0.3757E+00,0.3761E+00,0.3765E+00,
```

```
0.3769E+00,0.3773E+00,0.3778E+00,0.3782E+00,0.3786E+00,
    0.3790E+00,0.3794E+00,0.3798E+00,0.3802E+00,0.3807E+00,
#
    0.3811E+00,0.3815E+00,0.3819E+00,0.3823E+00,0.3827E+00,
#
    0.3831E+00,0.3835E+00,0.3839E+00,0.3844E+00,0.3848E+00/
#
    DATA (PCGTF2(I), I=451,500)/
    0.3852E+00,0.3856E+00,0.3860E+00,0.3864E+00,0.3868E+00,
#
    0.3872E+00,0.3876E+00,0.3880E+00,0.3884E+00,0.3889E+00,
#
    0.3893E+00,0.3897E+00,0.3901E+00,0.3905E+00,0.3909E+00,
#
    0.3913E+00,0.3917E+00,0.3921E+00,0.3925E+00,0.3930E+00,
#
    0.3934E+00,0.3938E+00,0.3942E+00,0.3946E+00,0.3950E+00,
#
    0.3954E+00,0.3958E+00,0.3963E+00,0.3967E+00,0.3971E+00,
#
    0.3975E+00,0.3979E+00,0.3983E+00,0.3987E+00,0.3992E+00,
#
    0.3996E+00,0.4000E+00,0.4004E+00,0.4008E+00,0.4013E+00,
#
    0.4017E+00.0.4021E+00,0.4025E+00,0.4030E+00,0.4034E+00,
    0.4038E+00,0.4042E+00,0.4047E+00,0.4051E+00,0.4055E+00/
    DATA (PCGTF2(I), I=501,550)/
    0.4060E+00,0.4063E+00,0.4066E+00,0.4069E+00,0.4072E+00,
#
    0.4075E+00,0.4078E+00,0.4081E+00,0.4084E+00,0.4087E+00,
#
    0.4090E+00,0.4093E+00,0.4096E+00,0.4100E+00,0.4103E+00,
#
    0.4106E+00,0.4109E+00,0.4112E+00,0.4115E+00,0.4118E+00,
#
    0.4121E+00,0.4124E+00,0.4127E+00,0.4131E+00,0.4134E+00,
#
    0.4137E+00,0.4140E+00,0.4143E+00,0.4146E+00,0.4149E+00,
#
    0.4152E+00,0.4156E+00,0.4159E+00,0.4162E+00,0.4165E+00,
#
    0.4168E+00,0.4171E+00,0.4174E+00,0.4178E+00,0.4181E+00,
#
    0.4184E+00,0.4187E+00,0.4190E+00,0.4193E+00,0.4197E+00,
#
    0.4200E+00,0.4203E+00,0.4206E+00,0.4209E+00,0.4212E+00/
    DATA (PCGTF2(I), I=551,600)/
    0.4216E+00,0.4219E+00,0.4222E+00,0.4225E+00,0.4228E+00,
#
    0.4231E+00,0.4235E+00,0.4238E+00,0.4241E+00,0.4244E+00,
#
    0.4247E+00,0.4251E+00,0.4254E+00,0.4257E+00,0.4260E+00,
#
    0.4264E+00,0.4267E+00,0.4270E+00,0.4273E+00,0.4276E+00,
#
    0.4280E+00,0.4283E+00,0.4286E+00,0.4289E+00,0.4293E+00,
#
    0.4296E+00,0.4299E+00,0.4302E+00,0.4306E+00,0.4309E+00,
#
    0.4312E+00,0.4315E+00,0.4319E+00,0.4322E+00,0.4325E+00,
#
    0.4328E+00,0.4332E+00,0.4335E+00,0.4338E+00,0.4341E+00,
#
    0.4345E+00,0.4348E+00,0.4351E+00,0.4355E+00,0.4358E+00,
#
    0.4361E+00,0.4364E+00,0.4368E+00,0.4371E+00,0.4374E+00/
#
    DATA (PCGTF2(I), I=601,650)/
    0.4378E+00,0.4381E+00,0.4384E+00,0.4388E+00,0.4391E+00,
#
    0.4394E+00,0.4397E+00,0.4401E+00,0.4404E+00,0.4407E+00,
#
    0.4411E+00,0.4414E+00,0.4417E+00,0.4421E+00,0.4424E+00,
#
    0.4427E+00,0.4431E+00,0.4434E+00,0.4437E+00,0.4441E+00,
#
    0.4444E+00,0.4447E+00,0.4451E+00,0.4454E+00,0.4458E+00,
#
    0.4461E+00,0.4464E+00,0.4468E+00,0.4471E+00,0.4474E+00,
#
    0.4478E+00,0.4481E+00,0.4485E+00,0.4488E+00,0.4491E+00,
#
    0.4495E+00,0.4498E+00,0.4502E+00,0.4505E+00,0.4508E+00,
    0.4512E+00,0.4515E+00,0.4519E+00,0.4522E+00,0.4525E+00,
#
    0.4529E+00,0.4532E+00,0.4536E+00,0.4539E+00,0.4542E+00/
#
    DATA (PCGTF2(I), I=651,700)/
    0.4546E+00,0.4549E+00,0.4553E+00,0.4556E+00,0.4560E+00,
#
    0.4563E+00,0.4566E+00,0.4570E+00,0.4573E+00,0.4577E+00,
#
#
    0.4580E+00,0.4584E+00,0.4587E+00,0.4591E+00,0.4594E+00,
    0.4598E+00,0.4601E+00,0.4605E+00,0.4608E+00,0.4611E+00,
#
    0.4615E+00,0.4618E+00,0.4622E+00,0.4625E+00,0.4629E+00,
#
    0.4632E+00,0.4636E+00,0.4639E+00,0.4643E+00,0.4646E+00,
#
#
    0.4650E+00,0.4653E+00,0.4657E+00,0.4660E+00,0.4664E+00,
    0.4667E+00,0.4671E+00,0.4675E+00,0.4678E+00,0.4682E+00,
    0.4685E+00,0.4689E+00,0.4692E+00,0.4696E+00,0.4699E+00,
    0.4703E+00,0.4706E+00,0.4710E+00,0.4713E+00,0.4717E+00/
    DATA (PCGTF2(I), I=701,750)/
```

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0.4721E+00,0.4724E+00,0.4728E+00,0.4731E+00,0.4735E+00,
    0.4738E+00,0.4742E+00,0.4746E+00,0.4749E+00,0.4753E+00,
#
    0.4756E+00,0.4760E+00,0.4764E+00,0.4767E+00,0.4771E+00,
    0.4774E+00,0.4778E+00,0.4782E+00,0.4785E+00,0.4789E+00,
#
    0.4792E+00,0.4796E+00,0.4800E+00,0.4803E+00,0.4807E+00,
    0.4810E+00,0.4814E+00,0.4818E+00,0.4821E+00,0.4825E+00,
#
    0.4829E+00,0.4832E+00,0.4836E+00,0.4840E+00,0.4843E+00,
    0.4847E+00,0.4851E+00,0.4854E+00,0.4858E+00,0.4862E+00,
#
    0.4865E+00,0.4869E+00,0.4873E+00,0.4876E+00,0.4880E+00,
    0.4884E+00,0.4887E+00,0.4891E+00,0.4895E+00,0.4898E+00/
#
    DATA (PCGTF2(I), I=751,800)/
    0.4902E+00,0.4906E+00,0.4909E+00,0.4913E+00,0.4917E+00,
#
    0.4921E+00,0.4924E+00,0.4928E+00,0.4932E+00,0.4935E+00,
#
    0.4939E+00,0.4943E+00,0.4947E+00,0.4950E+00,0.4954E+00,
    0.4958E+00,0.4962E+00,0.4965E+00,0.4969E+00,0.4973E+00,
#
    0.4977E+00,0.4980E+00,0.4984E+00,0.4988E+00,0.4992E+00,
#
    0.4995E+00,0.4999E+00,0.5003E+00,0.5007E+00,0.5010E+00,
#
    0.5014E+00,0.5018E+00,0.5022E+00,0.5026E+00,0.5029E+00,
#
    0.5033E+00,0.5037E+00,0.5041E+00,0.5045E+00,0.5048E+00,
#
    0.5052E+00,0.5056E+00,0.5060E+00,0.5064E+00,0.5068E+00,
#
    0.5071E+00,0.5075E+00,0.5079E+00,0.5083E+00,0.5087E+00/
    DATA (PCGTF2(I), I=801,850)/
    0.5091E+00,0.5094E+00,0.5098E+00,0.5102E+00,0.5106E+00,
#
    0.5110E+00,0.5114E+00,0.5117E+00,0.5121E+00,0.5125E+00,
#
    0.5129E+00,0.5133E+00,0.5137E+00,0.5141E+00,0.5145E+00,
#
    0.5148E+00,0.5152E+00,0.5156E+00,0.5160E+00,0.5164E+00,
#
    0.5168E+00,0.5172E+00,0.5176E+00,0.5180E+00,0.5184E+00,
#
    0.5187E+00,0.5191E+00,0.5195E+00,0.5199E+00,0.5203E+00,
#
    0.5207E+00,0.5211E+00,0.5215E+00,0.5219E+00,0.5223E+00,
#
    0.5227E+00,0.5231E+00,0.5235E+00,0.5239E+00,0.5243E+00,
#
    0.5246E+00,0.5250E+00,0.5254E+00,0.5258E+00,0.5262E+00,
#
    0.5266E+00,0.5270E+00,0.5274E+00,0.5278E+00,0.5282E+00/
#
    DATA (PCGTF2(I), I=851,900)/
    0.5286E+00,0.5290E+00,0.5294E+00,0.5298E+00,0.5302E+00,
#
    0.5306E+00,0.5310E+00,0.5314E+00,0.5318E+00,0.5322E+00,
#
    0.5326E+00,0.5330E+00,0.5334E+00,0.5338E+00,0.5342E+00,
#
    0.5346E+00,0.5350E+00,0.5354E+00,0.5358E+00,0.5363E+00,
#
     0.5367E+00,0.5371E+00,0.5375E+00,0.5379E+00,0.5383E+00,
#
     0.5387E+00,0.5391E+00,0.5395E+00,0.5399E+00,0.5403E+00,
 #
     0.5407E+00,0.5411E+00,0.5415E+00,0.5419E+00,0.5424E+00,
 #
     0.5428E+00,0.5432E+00,0.5436E+00,0.5440E+00,0.5444E+00,
 #
     0.5448E+00,0.5452E+00,0.5456E+00,0.5460E+00,0.5465E+00,
 #
     0.5469E+00,0.5473E+00,0.5477E+00,0.5481E+00,0.5485E+00/
 #
     DATA (PCGTF2(I), I=901,950)/
     0.5489E+00,0.5494E+00,0.5498E+00,0.5502E+00,0.5506E+00,
 #
     0.5510E+00,0.5514E+00,0.5518E+00,0.5523E+00,0.5527E+00,
 #
     0.5531E+00,0.5535E+00,0.5539E+00,0.5543E+00,0.5548E+00,
     0.5552E+00,0.5556E+00,0.5560E+00,0.5564E+00,0.5569E+00,
 #
     0.5573E+00,0.5577E+00,0.5581E+00,0.5585E+00,0.5590E+00,
 #
     0.5594E+00,0.5598E+00,0.5602E+00,0.5607E+00,0.5611E+00,
 #
     0.5615E+00,0.5619E+00,0.5624E+00,0.5628E+00,0.5632E+00,
     0.5636E+00,0.5641E+00,0.5645E+00,0.5649E+00,0.5653E+00,
 #
     0.5658E+00,0.5662E+00,0.5666E+00,0.5670E+00,0.5675E+00,
 #
     0.5679E+00,0.5683E+00,0.5688E+00,0.5692E+00,0.5696E+00/
 #
     DATA (PCGTF2(I), I=951, Nrigs)/
     0.5700E+00,0.5705E+00,0.5709E+00,0.5713E+00,0.5718E+00,
 #
     0.5722E+00,0.5726E+00,0.5731E+00,0.5735E+00,0.5739E+00,
 #
     0.5744E+00,0.5748E+00,0.5752E+00,0.5757E+00,0.5761E+00,
     0.5765E+00,0.5770E+00,0.5774E+00,0.5778E+00,0.5783E+00,
     0.5787E+00,0.5791E+00,0.5796E+00,0.5800E+00,0.5805E+00,
```

```
0.5809E+00,0.5813E+00,0.5818E+00,0.5822E+00,0.5826E+00,
#
    0.5831E+00,0.5835E+00,0.5840E+00,0.5844E+00,0.5849E+00,
#
    0.5853E+00,0.5857E+00,0.5862E+00,0.5866E+00,0.5871E+00,
#
    0.5875E+00,0.5879E+00,0.5884E+00,0.5888E+00,0.5893E+00,
#
    0.5897E+00,0.5902E+00,0.5906E+00,0.5911E+00,0.5915E+00,
#
#
    0.5920E+00/
    DATA (PCGTF3(I), I=1,50)/
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00,
#
    O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00,
#
    O.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00,
#
    O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    O.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00/
#
    DATA (PCGTF3(I), I=51,100)/
    O.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    O.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    O.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00,
#
    O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00,
#
    O.0000E+00,O.0000E+00,O.0000E+00,O.0000E+00,O.0000E+00/
#
    DATA (PCGTF3(I), I=101, 150)/
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    O.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00,
#
    O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.2036E-05,0.4190E-05,0.6583E-05,0.9331E-05,
#
    0.1256E-04,0.1637E-04,0.2091E-04,0.2627E-04,0.3258E-04/
#
    DATA (PCGTF3(I), I=151, 200)/
    0.3997E-04,0.4852E-04,0.5826E-04,0.6919E-04,0.8129E-04,
#
#
    0.9459E-04,0.1091E-03,0.1247E-03,0.1415E-03,0.1595E-03,
#
    0.1787E-03,0.1991E-03,0.2206E-03,0.2433E-03,0.2673E-03,
    0.2928E-03,0.3200E-03,0.3492E-03,0.3805E-03,0.4142E-03,
#
    0.4506E-03,0.4898E-03,0.5321E-03,0.5776E-03,0.6267E-03,
#
#
    0.6795E-03,0.7362E-03,0.7973E-03,0.8631E-03,0.9339E-03,
    0.1010E-02,0.1092E-02,0.1180E-02,0.1275E-02,0.1377E-02,
#
    0.1486E-02,0.1602E-02,0.1726E-02,0.1859E-02,0.2001E-02,
#
    0.2151E-02,0.2311E-02,0.2481E-02,0.2661E-02,0.2852E-02,
#
    0.3053E-02,0.3266E-02,0.3490E-02,0.3726E-02,0.3975E-02/
#
    DATA (PCGTF3(I), I=201, 250)/
    0.4237E-02,0.4511E-02,0.4798E-02,0.5097E-02,0.5406E-02,
#
    0.5727E-02,0.6057E-02,0.6397E-02,0.6746E-02,0.7102E-02,
#
    0.7466E-02,0.7837E-02,0.8214E-02,0.8597E-02,0.8984E-02,
#
    0.9376E-02,0.9772E-02,0.1017E-01,0.1057E-01,0.1097E-01,
#
    0.1138E-01,0.1178E-01,0.1219E-01,0.1259E-01,0.1299E-01,
#
    0.1339E-01,0.1379E-01,0.1419E-01,0.1458E-01,0.1498E-01,
#
    0.1537E-01,0.1577E-01,0.1617E-01,0.1657E-01,0.1697E-01,
#
#
    0.1738E-01,0.1779E-01,0.1821E-01,0.1864E-01,0.1907E-01,
#
    0.1951E-01,0.1996E-01,0.2041E-01,0.2088E-01,0.2136E-01,
```

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0.2185E-01,0.2235E-01,0.2287E-01,0.2340E-01,0.2394E-01/
    DATA (PCGTF3(I), I=251,300)/
    0.2450E-01,0.2508E-01,0.2567E-01,0.2628E-01,0.2690E-01,
#
    0.2753E-01,0.2818E-01,0.2884E-01,0.2951E-01,0.3020E-01,
    0.3089E-01,0.3160E-01,0.3231E-01,0.3304E-01,0.3377E-01,
#
    0.3452E-01,0.3527E-01,0.3602E-01,0.3679E-01,0.3756E-01,
#
    0.3834E-01,0.3912E-01,0.3990E-01,0.4069E-01,0.4149E-01,
#
    0.4228E-01,0.4308E-01,0.4388E-01,0.4468E-01,0.4548E-01,
#
    0.4629E-01,0.4709E-01,0.4789E-01,0.4869E-01,0.4949E-01,
#
    0.5028E-01,0.5107E-01,0.5186E-01,0.5264E-01,0.5342E-01,
#
    0.5419E-01,0.5496E-01,0.5572E-01,0.5648E-01,0.5722E-01,
#
    0.5796E-01,0.5869E-01,0.5941E-01,0.6012E-01,0.6082E-01/
#
    DATA (PCGTF3(I), I=301,350)/
    0.6151E-01,0.6219E-01,0.6285E-01,0.6351E-01,0.6416E-01,
#
    0.6479E-01,0.6542E-01,0.6604E-01,0.6665E-01,0.6726E-01,
#
    0.6785E-01,0.6844E-01,0.6902E-01,0.6960E-01,0.7017E-01,
#
    0.7074E-01,0.7130E-01,0.7186E-01,0.7241E-01,0.7296E-01,
#
    0.7351E-01,0.7406E-01,0.7460E-01,0.7515E-01,0.7569E-01,
#
    0.7623E-01,0.7677E-01,0.7732E-01,0.7786E-01,0.7841E-01,
#
    0.7896E-01,0.7951E-01,0.8006E-01,0.8062E-01,0.8118E-01,
#
    0.8174E-01,0.8231E-01,0.8289E-01,0.8347E-01,0.8405E-01,
    0.8465E-01,0.8525E-01,0.8586E-01,0.8647E-01,0.8710E-01,
#
    0.8773E-01,0.8837E-01,0.8902E-01,0.8969E-01,0.9036E-01/
#
    DATA (PCGTF3(I), I=351,400)/
    0.9105E-01,0.9174E-01,0.9245E-01,0.9317E-01,0.9390E-01,
#
    0.9464E-01,0.9540E-01,0.9616E-01,0.9693E-01,0.9772E-01,
#
    0.9851E-01,0.9931E-01,0.1001E+00,0.1009E+00,0.1018E+00,
#
    0.1026E+00,0.1035E+00,0.1043E+00,0.1052E+00,0.1061E+00,
#
    0.1070E+00,0.1078E+00,0.1087E+00,0.1097E+00,0.1106E+00,
#
    0.1115E+00,0.1124E+00,0.1133E+00,0.1143E+00,0.1152E+00,
#
    0.1162E+00,0.1171E+00,0.1181E+00,0.1191E+00,0.1201E+00,
#
    0.1210E+00,0.1220E+00,0.1230E+00,0.1240E+00,0.1250E+00,
#
    0.1260E+00,0.1270E+00,0.1280E+00,0.1290E+00,0.1301E+00,
#
    0.1311E+00,0.1321E+00,0.1331E+00,0.1342E+00,0.1352E+00/
#
    DATA (PCGTF3(I), I=401,450)/
    0.1362E+00,0.1373E+00,0.1383E+00,0.1394E+00,0.1404E+00,
#
    0.1414E+00,0.1425E+00,0.1435E+00,0.1446E+00,0.1456E+00,
#
    0.1467E+00,0.1478E+00,0.1488E+00,0.1499E+00,0.1509E+00,
#
    0.1520E+00,0.1530E+00,0.1541E+00,0.1552E+00,0.1562E+00,
#
    0.1573E+00,0.1584E+00,0.1594E+00,0.1605E+00,0.1616E+00,
#
     0.1626E+00,0.1637E+00,0.1648E+00,0.1658E+00,0.1669E+00,
#
     0.1680E+00,0.1690E+00,0.1701E+00,0.1712E+00,0.1722E+00,
#
     0.1733E+00,0.1744E+00,0.1754E+00,0.1765E+00,0.1776E+00,
#
     0.1786E+00,0.1797E+00,0.1807E+00,0.1818E+00,0.1829E+00,
 #
     0.1839E+00,0.1850E+00,0.1860E+00,0.1871E+00,0.1882E+00/
#
     DATA (PCGTF3(I), I=451,500)/
     0.1892E+00,0.1903E+00,0.1913E+00,0.1924E+00,0.1934E+00,
 #
     0.1945E+00,0.1955E+00,0.1965E+00,0.1976E+00,0.1986E+00,
 #
     0.1997E+00,0.2007E+00,0.2017E+00,0.2027E+00,0.2038E+00,
 #
     0.2048E+00,0.2058E+00,0.2068E+00,0.2079E+00,0.2089E+00,
 #
     0.2099E+00,0.2109E+00,0.2119E+00,0.2129E+00,0.2139E+00,
 #
     0.2149E+00,0.2159E+00,0.2169E+00,0.2179E+00,0.2189E+00,
 #
     0.2198E+00,0.2208E+00,0.2218E+00,0.2227E+00,0.2237E+00,
 #
     0.2247E+00,0.2256E+00,0.2266E+00,0.2275E+00,0.2285E+00,
 #
     0.2294E+00,0.2304E+00,0.2313E+00,0.2322E+00,0.2331E+00,
 #
     0.2341E+00,0.2350E+00,0.2359E+00,0.2368E+00,0.2377E+00/
     DATA (PCGTF3(I), I=501,550)/
     0.2386E+00,0.2395E+00,0.2404E+00,0.2412E+00,0.2421E+00,
 #
     0.2430E+00,0.2439E+00,0.2447E+00,0.2456E+00,0.2464E+00,
 #
     0.2473E+00,0.2481E+00,0.2490E+00,0.2498E+00,0.2506E+00,
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0.2515E+00,0.2523E+00,0.2531E+00,0.2539E+00,0.2547E+00,
   0.2555E+00,0.2563E+00,0.2571E+00,0.2579E+00,0.2587E+00,
#
   0.2595E+00,0.2603E+00,0.2611E+00,0.2619E+00,0.2626E+00,
#
    0.2634E+00,0.2642E+00,0.2649E+00,0.2657E+00,0.2664E+00,
#
    0.2672E+00,0.2680E+00,0.2687E+00,0.2694E+00,0.2702E+00,
#
    0.2709E+00,0.2717E+00,0.2724E+00,0.2731E+00,0.2739E+00,
#
    0.2746E+00,0.2753E+00,0.2760E+00,0.2768E+00,0.2775E+00/
#
   DATA (PCGTF3(I), I=551,600)/
    0.2782E+00,0.2789E+00,0.2796E+00,0.2803E+00,0.281CE+00,
#
    0.2817E+00,0.2824E+00,0.2831E+00,0.2838E+00,0.2845E+00,
#
    0.2852E+00,0.2859E+00,0.2866E+00,0.2873E+00,0.2880E+00,
#
    0.2887E+00,0.2894E+00,0.2900E+00,0.2907E+00,0.2914E+00,
#
    0.2921E+00,0.2928E+00,0.2934E+00,0.2941E+00,0.2948E+00,
#
    0.2955E+00,0.2961E+00,0.2968E+00,0.2975E+00,0.2982E+00,
#
#
    0.2988E+00,0.2995E+00,0.3002E+00,0.3008E+00,0.3015E+00,
    0.3022E+00,0.3028E+00,0.3035E+00,0.3042E+00,0.3048E+00,
#
    0.3055E+00,0.3062E+00,0.3069E+00,0.3075E+00,0.3082E+00,
#
#
    0.3089E+00,0.3095E+00,0.3102E+00,0.3109E+00,0.3115E+00/
    DATA (PCGTF3(I), I=601,650)/
    0.3122E+00,0.3129E+00,0.3135E+00,0.3142E+00,0.3149E+00,
#
    0.3156E+00,0.3162E+00,0.3169E+00,0.3176E+00,0.3182E+00,
#
#
    0.3189E+00,0.3196E+00,0.3203E+00,0.3209E+00,0.3216E+00,
    0.3223E+00,0.3230E+00,0.3237E+00,0.3243E+00,0.325CE+00,
#
    0.3257E+00,0.3264E+00,0.3270E+00,0.3277E+00,0.3284E+00,
#
    0.3291E+00.0.3298E+00.0.3304E+00.0.3311E+00.0.3318E+00.
#
#
    0.3325E+00,0.3332E+00,0.3339E+00,0.3345E+00,0.3352E+00,
    0.3359E+00,0.3366E+00,0.3373E+00,0.3379E+00,0.3386E+00,
#
    0.3393E+00,0.3400E+00,0.3407E+00,0.3414E+00,0.3421E+00,
#
    0.3427E+00,0.3434E+00,0.3441E+00,0.3448E+00,0.3455E+00/
#
    DATA (PCGTF3(I), I=651,700)/
    0.3462E+00,0.3469E+00,0.3475E+00,0.3482E+00,0.3489E+00,
#
    0.3496E+00,0.3503E+00,0.3510E+00,0.3517E+00,0.3523E+00,
#
    0.3530E+00,0.3537E+00,0.3544E+00,0.3551E+00,0.3558E+00,
#
    0.3565E+00,0.3572E+00,0.3578E+00,0.3585E+00,0.3592E+00,
#
    0.3599E+00,0.3606E+00,0.3613E+00,0.3620E+00,0.3627E+00,
#
    0.3634E+00,0.3640E+00,0.3647E+00,0.3654E+00,0.3661E+00,
#
    0.3668E+00,0.3675E+00,0.3682E+00,0.3689E+00,0.3695E+00,
#
#
    0.3702E+00,0.3709E+00,0.3716E+00,0.3723E+00,0.3730E+00,
    0.3737E+00,0.3743E+00,0.3750E+00,0.3757E+00,0.3764E+00,
#
    0.3771E+00,0.3778E+00,0.3785E+00,0.3792E+00,0.3798E+00/
#
    DATA (PCGTF3(I), I=701,750)/
    0.3805E+00,0.3812E+00,0.3819E+00,0.3826E+00,0.3833E+00,
#
#
    0.3839E+00,0.3846E+00,0.3853E+00,0.3860E+00,0.3867E+00,
#
    0.3874E+00,0.3880E+00,0.3887E+00,0.3894E+00,0.3901E+00,
    0.3908E+00,0.3915E+00,0.3921E+00,0.3928E+00,0.3935E+00,
    0.3942E+00,0.3949E+00,0.3955E+00,0.3962E+00,0.3969E+00,
#
#
    0.3976E+00,0.3983E+00,0.3989E+00,0.3996E+00,0.4003E+00,
#
    0.4010E+00,0.4016E+00,0.4023E+00,0.4030E+00,0.4037E+00,
#
    0.4044E+00,0.4050E+00,0.4057E+00,0.4064E+00,0.4070E+00,
#
    0.4077E+00,0.4084E+00,0.4091E+00,0.4097E+00,0.4104E+00,
#
    0.4111E+00,0.4117E+00,0.4124E+00,0.4131E+00,0.4138E+00/
    DATA (PCGTF3(I), I=751,800)/
    0.4144E+00,0.4151E+00,0.4158E+00,0.4164E+00,0.4171E+00,
#
#
    0.4178E+00,0.4184E+00,0.4191E+00,0.4197E+00,0.4204E+00,
#
    0.4211E+00,0.4217E+00,0.4224E+00,0.4231E+00,0.4237E+00,
#
    0.4244E+00,0.4250E+00,0.4257E+00,0.4264E+00,0.4270E+00,
    0.4277E+00,0.4283E+00,0.4290E+00,0.4296E+00,0.4303E+00,
    0.4309E+00,0.4316E+00,0.4322E+00,0.4329E+00,0.4335E+00,
#
#
    0.4342E+00,0.4348E+00,0.4355E+00,0.4361E+00,0.4368E+00,
    0.4374E+00,0.4381E+00,0.4387E+00,0.4394E+00,0.4400E+00,
```

```
0.4406E+00,0.4413E+00,0.4419E+00,0.4426E+00,0.4432E+00,
#
    0.4438E+00,0.4445E+00,0.4451E+00,0.4458E+00,0.4464E+00/
    DATA (PCGTF3(I), I=801,850)/
    0.4470E+00,0.4474E+00,0.4478E+00,0.4481E+00,0.4485E+00,
#
    0.4489E+00,0.4493E+00,0.4497E+00,0.4500E+00,0.4504E+00,
#
    0.4508E+00,0.4512E+00,0.4516E+00,0.4519E+00,0.4523E+00,
    0.4527E+00,0.4531E+00,0.4535E+00,0.4538E+00,0.4542E+00,
    0.4546E+00,0.4550E+00,0.4554E+00,0.4558E+00,0.4561E+00,
    0.4565E+00,0.4569E+00,0.4573E+00,0.4577E+00,0.4581E+00,
#
#
    0.4584E+00,0.4588E+00,0.4592E+00,0.4596E+00,0.4600E+00,
#
    0.4604E+00,0.4608E+00,0.4612E+00,0.4615E+00,0.4619E+00,
#
    0.4623E+00,0.4627E+00,0.4631E+00,0.4635E+00,0.4639E+00,
#
    0.4643E+00,0.4647E+00,0.4650E+00,0.4654E+00,0.4658E+00/
    DATA (PCGTF3(I), I=851, 900)/
    0.4662E+00,0.4666E+00,0.4670E+00,0.4674E+00,0.4678E+00,
    0.4682E+00,0.4686E+00,0.4690E+00,0.4694E+00,0.4698E+00,
#
#
    0.4702E+00,0.4706E+00,0.4710E+00,0.4713E+00,0.4717E+00,
    0.4721E+00,0.4725E+00,0.4729E+00,0.4733E+00,0.4737E+00,
    0.4741E+00,0.4745E+00,0.4749E+00,0.4753E+00,0.4757E+00,
#
    0.4761E+00,0.4765E+00,0.4769E+00,0.4773E+00,0.4777E+00,
#
    0.4781E+00,0.4785E+00,0.4789E+00,0.4793E+00,0.4798E+00,
    0.4802E+00,0.4806E+00,0.4810E+00,0.4814E+00,0.4818E+00,
    0.4822E+00,0.4826E+00,0.4830E+00,0.4834E+00,0.4838E+00,
#
    0.4842E+00,0.4846E+00,0.4850E+00,0.4854E+00,0.4858E+00/
    DATA (PCGTF3(I), I=901, 950)/
#
    0.4863E+00,0.4867E+00,0.4871E+00,0.4875E+00,0.4879E+00,
#
    0.4883E+00,0.4887E+00,0.4891E+00,0.4895E+00,0.4899E+00,
#
    0.4904E+00,0.4908E+00,0.4912E+00,0.4916E+00,0.4920E+00,
    0.4924E+00,0.4928E+00,0.4933E+00,0.4937E+00,0.4941E+00,
    0.4945E+00,0.4949E+00,0.4953E+00,0.4958E+00,0.4962E+00,
#
#
    0.4966E+00,0.4970E+00,0.4974E+00,0.4978E+00,0.4983E+00,
#
    0.4987E+00,0.4991E+00,0.4995E+00,0.4999E+00,0.5004E+00,
    0.5008E+00,0.5012E+00,0.5016E+00,0.5020E+00,0.5025E+00,
    0.5029E+00,0.5033E+00,0.5037E+00,0.5042E+00,0.5046E+00,
#
    0.5050E+00,0.5054E+00,0.5059E+00,0.5063E+00,0.5067E+00/
    DATA (PCGTF3(I), I=951, Nrigs)/
    0.5071E+00,0.5076E+00,0.5080E+00,0.5084E+00,0.5088E+00,
#
    0.5093E+00,0.5097E+00,0.5101E+00,0.5106E+00,0.5110E+00,
#
    0.5114E+00,0.5119E+00,0.5123E+00,0.5127E+00,0.5131E+00,
#
    0.5136E+00,0.5140E+00,0.5144E+00,0.5149E+00,0.5153E+00,
    0.5157E+00,0.5162E+00,0.5166E+00,0.5170E+00,0.5175E+00,
    0.5179E+00,0.5184E+00,0.5188E+00,0.5192E+00,0.5197E+00,
    0.5201E+00,0.5205E+00,0.5210E+00,0.5214E+00,0.5219E+00,
    0.5223E+00,0.5227E+00,0.5232E+00,0.5236E+00,0.5241E+00,
    0.5245E+00,0.5249E+00,0.5254E+00,0.5258E+00,0.5263E+00,
#
    0.5267E+00,0.5271E+00,0.5276E+00,0.5280E+00,0.5285E+00,
    0.5289E+00/
    DATA (PCGTF4(I), I=1,50)/
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00/
    DATA (PCGTF4(I), I=51,100)/
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
```

```
0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
    O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00, O.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00/
#
    DATA (PCGTF4(I), I=101, 150)/
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,
#
    0.0000E+00,0.0000E+00,0.0000E+00,0.0000E+00,0.1998E-04,
#
    0.3997E-04,0.5863E-04,0.7762E-04,0.9727E-04,0.1179E-03,
#
    0.1399E-03,0.1636E-03,0.1896E-03,0.2185E-03,0.2510E-03,
#
    0.2878E-03,0.3293E-03,0.3757E-03,0.4268E-03,0.4827E-03,
#
    0.5433E-03,0.6086E-03,0.6784E-03,0.7529E-03,0.8318E-03/
   DATA (PCGTF4(I), I=151, 200)/
#
    0.9153E-03,0.1003E-02,0.1096E-02,0.1195E-02,0.1299E-02,
#
    0.1410E-02,0.1528E-02,0.1653E-02,0.1787E-02,0.1929E-02,
#
    0.2081E-02,0.2242E-02,0.2413E-02,0.2595E-02,0.2788E-02,
#
    0.2991E-02,0.3205E-02,0.3428E-02,0.3660E-02,0.3901E-02,
#
    0.4151E-02,0.4409E-02,0.4675E-02,0.4948E-02,0.5229E-02,
#
    0.5516E-02,0.5809E-02,0.6109E-02,0.6416E-02,0.6730E-02,
#
    0.7052E-02,0.7381E-02,0.7719E-02,0.8064E-02,0.8419E-02,
#
    0.8782E-02,0.9154E-02,0.9536E-02,0.9928E-02,0.1033E-01,
    0.1074E-01,0.1116E-01,0.1160E-01,0.1204E-01,0.1250E-01,
#
#
    0.1296E-01,0.1344E-01,0.1394E-01,0.1444E-01,0.1496E-01/
   DATA (PCGTF4(I), I=201,250)/
#
    0.1549E-01,0.1603E-01,0.1659E-01,0.1715E-01,0.1773E-01,
    0.1832E-01,0.1892E-01,0.1952E-01,0.2013E-01,0.2074E-01,
#
    0.2136E-01,0.2198E-01,0.2261E-01,0.2323E-01,0.2386E-01,
#
    0.2448E-01,0.2510E-01,0.2572E-01,0.2634E-01,0.2695E-01,
#
    0.2755E-01,0.2815E-01,0.2874E-01,0.2932E-01,0.2990E-01,
#
    0.3046E-01,0.3100E-01,0.3154E-01,0.3207E-01,0.3259E-01,
#
    0.3310E-01,0.3360E-01,0.3410E-01,0.3459E-01,0.3508E-01,
#
    0.3556E-01,0.3604E-01,0.3652E-01,0.3700E-01,0.3748E-01,
#
    0.3796E-01,0.3844E-01,0.3893E-01,0.3942E-01,0.3992E-01,
#
    0.4042E-01,0.4093E-01,0.4145E-01,0.4197E-01,0.4251E-01/
   DATA (PCGTF4(I), I=251, 300)/
#
    0.4306E-01,0.4362E-01,0.4419E-01,0.4477E-01,0.4536E-01,
    0.4596E-01,0.4658E-01,0.4720E-01,0.4783E-01,0.4847E-01,
#
#
    0.4912E-01,0.4978E-01,0.5044E-01,0.5111E-01,0.5179E-01,
#
    0.5248E-01,0.5317E-01,0.5386E-01,0.5457E-01,0.5527E-01,
#
    0.5598E-01,0.5670E-01,0.5742E-01,0.5814E-01,0.5886E-01,
#
    0.5959E-01,0.6032E-01,0.6105E-01,0.6178E-01,0.6251E-01,
#
    0.6324E-01,0.6397E-01,0.6470E-01,0.6543E-01,0.6616E-01,
#
    0.6689E-01,0.6762E-01,0.6834E-01,0.6906E-01,0.6978E-01,
    0.7049E-01,0.7120E-01,0.7191E-01,0.7261E-01,0.7331E-01,
#
    0.7399E-01,0.7468E-01,0.7536E-01,0.7603E-01,0.7669E-01/
   DATA (PCGTF4(I), I=301,350)/
#
    0.7735E-01,0.7799E-01,0.7863E-01,0.7927E-01,0.7989E-01,
#
    0.8051E-01,0.8112E-01,0.8173E-01,0.8233E-01,0.8293E-01,
#
    0.8352E-01,0.8411E-01,0.8469E-01,0.8527E-01,0.8585E-01,
#
    0.8642E-01,0.8699E-01,0.8756E-01,0.8813E-01,0.8869E-01,
#
    0.8926E-01,0.8982E-01,0.9038E-01,0.9095E-01,0.9151E-01,
    0.9208E-01,0.9264E-01,0.9321E-01,0.9378E-01,0.9435E-01,
```

```
0.9492E-01,0.9550E-01,0.9607E-01,0.9666E-01,0.9724E-01,
     0.9784E-01,0.9843E-01,0.9903E-01,0.9964E-01,0.1003E+00,
 #
     0.1009E+00,0.1015E+00,0.1021E+00,0.1028E+00,0.1034E+00,
 #
     0.1041E+00,0.1047E+00,0.1054E+00,0.1061E+00,0.1068E+00/
     DATA (PCGTF4(I), I=351, 400)/
     0.1075E+00,0.1082E+00,0.1089E+00,0.1096E+00,0.1104E+00,
 #
     0.1111E+00,0.1119E+00,0.1126E+00,0.1134E+00,0.1142E+00,
     0.1150E+00,0.1158E+00,0.1166E+00,0.1174E+00,0.1182E+00,
 #
     0.1191E+00,0.1199E+00,0.1208E+00,0.1216E+00,0.1225E+00,
 #
     0.1233E+00,0.1242E+00,0.1251E+00,0.1260E+00,0.1268E+00,
 #
    0.1277E+00,0.1286E+00,0.1295E+00,0.1304E+00,0.1314E+00,
    0.1323E+00,0.1332E+00,0.1341E+00,0.1350E+00,0.1360E+00,
    0.1369E+00,0.1379E+00,0.1388E+00,0.1397E+00,0.1407E+00,
    0.1416E+00,0.1426E+00,0.1435E+00,0.1445E+00,0.1454E+00,
    0.1464E+00,0.1474E+00,0.1483E+00,0.1493E+00,0.1502E+00/
#
    DATA (PCGTF4(I), I=401,450)/
#
    0.1512E+00,0.1522E+00,0.1531E+00,0.1541E+00,0.1550E+00,
#
    0.1560E+00,0.1570E+00,0.1579E+00,0.1589E+00,0.1598E+00.
#
    0.1608E+00,0.1618E+00,0.1627E+00,0.1637E+00,0.1647E+00,
#
    0.1656E+00,0.1666E+00,0.1675E+00,0.1685E+00,0.1695E+00,
    0.1704E+00,0.1714E+00,0.1723E+00,0.1733E+00,0.1743E+00,
#
    0.1752E+00,0.1762E+00,0.1771E+00,0.1781E+00,0.1791E+00,
    0.1800E+00,0.1810E+00,0.1819E+00,0.1829E+00,0.1838E+00,
    0.1848E+00,0.1858E+00,0.1867E+00,0.1877E+00,0.1886E+00,
#
    0.1896E+00,0.1905E+00,0.1915E+00,0.1925E+00,0.1934E+00,
#
    0.1944E+00,0.1953E+00,0.1963E+00,0.1972E+00,0.1982E+00/
    DATA (PCGTF4(I), I=451,500)/
#
    0.1991E+00,0.2001E+00,0.2010E+00,0.2020E+00,0.2029E+00,
#
    0.2039E+00,0.2048E+00,0.2058E+00,0.2067E+00,0.2077E+00,
    0.2086E+00,0.2096E+00,0.2105E+00,0.2115E+00,0.2124E+00,
    0.2134E+00,0.2143E+00,0.2153E+00,0.2162E+00,0.2172E+00,
    0.2181E+00,0.2191E+00,0.2200E+00,0.2209E+00,0.2219E+00,
#
#
    0.2228E+00,0.2238E+00,0.2247E+00,0.2256E+00,0.2266E+00,
    0.2275E+00,0.2285E+00,0.2294E+00,0.2303E+00,0.2313E+00,
#
    0.2322E+00,0.2331E+00,0.2341E+00,0.2350E+00,0.2359E+00,
    0.2369E+00,0.2378E+00,0.2387E+00,0.2396E+00,0.2406E+00,
    0.2415E+00,0.2424E+00,0.2434E+00,0.2443E+00,0.2452E+00/
    DATA (PCGTF4(I), I=501,550)/
#
    0.2461E+00,0.2471E+00,0.2480E+00,0.2489E+00,0.2498E+00,
#
    0.2507E+00,0.2517E+00,0.2526E+00,0.2535E+00,0.2544E+00.
#
    0.2553E+00,0.2562E+00,0.2571E+00,0.2581E+00,0.2590E+00,
    0.2599E+00,0.2608E+00,0.2617E+00,0.2626E+00,0.2635E+00,
    0.2644E+00,0.2653E+00,0.2662E+00,0.2671E+00,0.2680E+00,
    0.2689E+00,0.2698E+00,0.2707E+00,0.2716E+00,0.2725E+00,
    0.2734E+00,0.2743E+00,0.2752E+00,0.2760E+00,0.2769E+00,
    0.2778E+00,0.2787E+00,0.2796E+00,0.2805E+00,0.2813E+00,
    0.2822E+00,0.2831E+00,0.2840E+00,0.2848E+00,0.2857E+00,
#
    0.2866E+00,0.2875E+00,0.2883E+00,0.2892E+00,0.2901E+00/
    DATA (PCGTF4(I), I=551,600)/
#
    0.2909E+00,0.2918E+00,0.2926E+00,0.2935E+00,0.2944E+00,
#
    0.2952E+00,0.2961E+00,0.2969E+00,0.2978E+00,0.2986E+00,
    0.2995E+00,0.3003E+00,0.3012E+00,0.3020E+00,0.3028E+00,
#
   0.3037E+00,0.3045E+00,0.3053E+00,0.3062E+00,0.3070E+00,
   0.3078E+00,0.3087E+00,0.3095E+00,0.3103E+00,0.3111E+00,
   0.3120E+00,0.3128E+00,0.3136E+00,0.3144E+00,0.3152E+00,
#
#
   0.3160E+00,0.3168E+00,0.3176E+00,0.3184E+00,0.3192E+00,
#
   0.3200E+00,0.3208E+00,0.3216E+00,0.3224E+00,0.3232E+00,
   0.3240E+00,0.3248E+00,0.3256E+00,0.3263E+00,0.3271E+00,
   0.3279E+00,0.3287E+00,0.3294E+00,0.3302E+00,0.3310E+00/
   DATA (PCGTF4(I), I=601,650)/
```

```
#
    0.3318E+00,0.3325E+00,0.3333E+00,0.3340E+00,0.3348E+00,
#
    0.3355E+00,0.3363E+00,0.3370E+00,0.3378E+00,0.3385E+00,
    0.3393E+00,0.3400E+00,0.3408E+00,0.3415E+00,0.3422E+00,
#
    0.3430E+00,0.3437E+00,0.3444E+00,0.3451E+00,0.3459E+00,
    0.3466E+00,0.3473E+00,0.3480E+00,0.3487E+00,0.3494E+00,
#
    0.3502E+00,0.3509E+00,0.3516E+00,0.3523E+00,0.3530E+00,
#
    0.3537E+00,0.3544E+00,0.3551E+00,0.3558E+00,0.3565E+00,
    0.3572E+00,0.3578E+00,0.3585E+00,0.3592E+00,0.3599E+00,
    0.3606E+00,0.3613E+00,0.3619E+00,0.3626E+00,0.3633E+00,
    0.3640E+00,0.3646E+00,0.3653E+00,0.3660E+00,0.3666E+00/
    DATA (PCGTF4(I), I=651,700)/
#
    0.3673E+00,0.3680E+00,0.3686E+00,0.3693E+00,0.3699E+00,
    0.3706E+00,0.3712E+00,0.3719E+00,0.3725E+00,0.3732E+00,
    0.3738E+00,0.3745E+00,0.3751E+00,0.3758E+00,0.3764E+00,
    0.3771E+00,0.3777E+00,0.3783E+00,0.3790E+00,0.3796E+00,
#
    0.3802E+00,0.3809E+00,0.3815E+00,0.3821E+00,0.3827E+00,
    0.3834E+00,0.3840E+00,0.3846E+00,0.3852E+00,0.3858E+00,
#
    0.3865E+00,0.3871E+00,0.3877E+00,0.3883E+00,0.3889E+00,
#
    0.3895E+00,0.3901E+00,0.3907E+00,0.3914E+00,0.3920E+00,
    0.3926E+00,0.3932E+00,0.3938E+00,0.3944E+00,0.3950E+00,
#
    0.3956E+00,0.3962E+00,0.3968E+00,0.3974E+00,0.3980E+00/
    DATA (PCGTF4(I), I=701,750)/
#
    0.3985E+00,0.3991E+00,0.3997E+00,0.4003E+00,0.4009E+00,
    0.4015E+00,0.4021E+00,0.4027E+00,0.4032E+00,0.4038E+00,
#
    0.4044E+00,0.4050E+00,0.4056E+00,0.4062E+00,0.4067E+00,
#
    0.4073E+00,0.4079E+00,0.4085E+00,0.4090E+00,0.4096E+00,
#
    0.4102E+00,0.4108E+00,0.4113E+00,0.4119E+00,0.4125E+00,
#
    0.4130E+00,0.4136E+00,0.4142E+00,0.4147E+00,0.4153E+00,
    0.4159E+00,0.4164E+00,0.4170E+00,0.4176E+00,0.4181E+00,
#
    0.4187E+00,0.4193E+00,0.4198E+00,0.4204E+00,0.4209E+00,
#
    0.4215E+00,0.4221E+00,0.4226E+00,0.4232E+00,0.4237E+00,
#
    0.4243E+00,0.4248E+00,0.4254E+00,0.4259E+00,0.4265E+00/
    DATA (PCGTF4(I), I=751,800)/
#
    0.4270E+00,0.4276E+00,0.4282E+00,0.4287E+00,0.4293E+00,
#
    0.4298E+00,0.4304E+00,0.4309E+00,0.4315E+00,0.4320E+00,
#
    0.4326E+00,0.4331E+00,0.4337E+00,0.4342E+00,0.4348E+00,
#
    0.4353E+00,0.4358E+00,0.4364E+00,0.4369E+00,0.4375E+00,
#
    0.4380E+00,0.4386E+00,0.4391E+00,0.4397E+00,0.4402E+00,
#
    0.4408E+00,0.4413E+00,0.4419E+00,0.4424E+00,0.4429E+00,
#
    0.4435E+00,0.4440E+00,0.4446E+00,0.4451E+00,0.4457E+00,
#
    0.4462E+00,0.4467E+00,0.4473E+00,0.4478E+00,0.4484E+00,
#
    0.4489E+00,0.4495E+00,0.4500E+00,0.4506E+00,0.4511E+00,
#
    0.4516E+00,0.4522E+00,0.4527E+00,0.4533E+00,0.4538E+00/
    DATA (PCGTF4(I), I=801,850)/
    0.4544E+00,0.4547E+00,0.4551E+00,0.4554E+00,0.4558E+00,
#
#
    0.4561E+00,0.4565E+00,0.4568E+00,0.4571E+00,0.4575E+00,
    0.4578E+00,0.4582E+00,0.4585E+00,0.4589E+00,0.4592E+00,
#
    0.4596E+00,0.4599E+00,0.4603E+00,0.4606E+00,0.4610E+00,
#
    0.4614E+00,0.4617E+00,0.4621E+00,0.4624E+00,0.4628E+00,
#
    0.4631E+00,0.4635E+00,0.4638E+00,0.4642E+00,0.4645E+00,
#
    0.4649E+00,0.4652E+00,0.4656E+00,0.4659E+00,0.4663E+00,
    0.4667E+00,0.4670E+00,0.4674E+00,0.4677E+00,0.4681E+00,
#
    0.4684E+00,0.4688E+00,0.4692E+00,0.4695E+00,0.4699E+00,
    0.4702E+00,0.4706E+00,0.4709E+00,0.4713E+00,0.4717E+00/
    DATA (PCGTF4(I), I=851,900)/
#
    0.4720E+00,0.4724E+00,0.4727E+00,0.4731E+00,0.4735E+00,
    0.4738E+00,0.4742E+00,0.4746E+00,0.4749E+00,0.4753E+00,
#
#
    0.4756E+00,0.4760E+00,0.4764E+00,0.4767E+00,0.4771E+00,
    0.4775E+00,0.4778E+00,0.4782E+00,0.4785E+00,0.4789E+00,
    0.4793E+00,0.4796E+00,0.4800E+00,0.4804E+00,0.4807E+00,
```

END

```
0.4811E+00,0.4815E+00,0.4818E+00,0.4822E+00,0.4826E+00,
#
    0.4829E+00,0.4833E+00,0.4837E+00,0.4841E+00,0.4844E+00,
#
    0.4848E+00,0.4852E+00,0.4855E+00,0.4859E+00,0.4863E+00,
    0.4866E+00,0.4870E+00,0.4874E+00,0.4878E+00,0.4881E+00,
#
#
    0.4885E+00,0.4889E+00,0.4892E+00,0.4896E+00,0.4900E+00/
    DATA (PCGTF4(I), I=901, 950)/
#
    0.4904E+00,0.4907E+00,0.4911E+00,0.4915E+00,0.4919E+00.
#
    0.4922E+00,0.4926E+00,0.4930E+00,0.4934E+00,0.4937E+00,
    0.4941E+00,0.4945E+00,0.4949E+00,0.4953E+00,0.4956E+00,
#
#
    0.4960E+00,0.4964E+00,0.4968E+00,0.4971E+00,0.4975E+00,
#
    0.4979E+00,0.4983E+00,0.4987E+00,0.4990E+00,0.4994E+00,
#
    0.4998E+00,0.5002E+00,0.5006E+00,0.5009E+00,0.5013E+00,
#
    0.5017E+00,0.5021E+00,0.5025E+00,0.5029E+00,0.5032E+00,
#
    0.5036E+00,0.5040E+00,0.5044E+00,0.5048E+00,0.5052E+00,
    0.5056E+00,0.5059E+00,0.5063E+00,0.5067E+00,0.5071E+00,
#
    0.5075E+00,0.5079E+00,0.5083E+00,0.5086E+00,0.5090E+00/
    DATA (PCGTF4(I), I=951, Nrigs)/
#
    0.5094E+00,0.5098E+00,0.5102E+00,0.5106E+00,0.5110E+00,
#
    0.5114E+00,0.5118E+00,0.5121E+00,0.5125E+00,0.5129E+00,
#
    0.5133E+00,0.5137E+00,0.5141E+00,0.5145E+00,0.5149E+00,
#
    0.5153E+00,0.5157E+00,0.5161E+00,0.5165E+00,0.5169E+00,
    0.5172E+00,0.5176E+00,0.5180E+00,0.5184E+00,0.5188E+00,
#
    0.5192E+00,0.5196E+00,0.5200E+00,0.5204E+00,0.5208E+00,
    0.5212E+00,0.5216E+00,0.5220E+00,0.5224E+00,0.5228E+00,
#
#
    0.5232E+00,0.5236E+00,0.5240E+00,0.5244E+00,0.5248E+00,
#
    0.5252E+00,0.5256E+00,0.5260E+00,0.5264E+00,0.5268E+00,
#
    0.5272E+00,0.5276E+00,0.5280E+00,0.5284E+00,0.5288E+00,
    0.5292E+00/
    DO I=1, Nrigs
     RigBins(I)=RIGPC(I)
  RETURN
```

```
С
                              Calculate Universal Time and Local Time. Note that Zlon
                              and period are not both required. Used as a consistency check
          C
          C
                             here.
                             IMPLICIT NONE
                             REAL Time, UTtimeInit, UTtime, Zlon, Period, TimeLocal
                             REAL secsperday
                             PARAMETER (secsperday=86400.0)
                             Integer Ndays, Ndaysloc
                             REAL TimeLoc
                             UTtime=UTtimeInit+time
                            Ndays=INT(UTtime/secsperday)
                            UTtime=UTtime - Ndays*secsperday
                            TimeLocal=UTtime+secsperday*Zlon/360.0
                            IF (TimeLocal .GE. secsperday) TimeLocal=TimeLocal-secsperday
                            RETURN
                            END
The state of the s
                                                          _______
                            SUBROUTINE CutputTransFcn(RigBins, TransFunc, GtransFile, OrbIncl,
                    #
                                                              Apogee, Perigee, AscNodeLong, AscNodeDisp, PerigDisp,
#
                                                              Zenith, Azimuth, UTtimeInit, Stormy, Shadow,
#
                                                              PreCalcGTFs, IPreCalc, Year, XLbounds,
20
                                                              ILbins, IprogNo)
L.S
IMPLICIT NONE
M C
li C
                           Format of header changed by AJT 8-21-96.
                           File names for L-bin results changed from .BN* to .GT* by AJT 11-6-96
₽ C
       C
                           IF one L-bin specified with L_{\min} > 0, make file extension .GT1
                           instead of .GTF. 11-21-96, PRB.
       C
                           REAL OrbIncl, Apogee, Perigee, AscNodeLong, AscNodeDisp, PerigDisp
       С
                           These variables are fixed at present and thus not included
       С
                           in header output.
                          REAL Zenith, Azimuth, UTtimeInit
       С
                          Shadow is not checked in determining output, since all GTF
       C
                          calculations included in GEOMAG96 are omnidirectional averages.
                          LOGICAL Shadow, Stormy, PreCalcGTFs, StormyPreCalc
                          INTEGER IPreCalc
      C
                         Name of output file, thus not included in output file header.
                          CHARACTER*80 GtransFile, TempFile
```

INTEGER I, Nrigs, NLvals, L, ILbins

SUBROUTINE ConvertTime(time,UTtimeInit,UTtime,Zlon,Period,

TimeLocal)

```
PARAMETER (Nrigs=1001, NLvals=10)
            REAL TransFunc (Nrigs, NLvals), RigBins (Nrigs)
            REAL XLbounds (NLvals), XLinfinite, Year
            PARAMETER (XLinfinite=1.0E+06)
            INTEGER ICREME96vno, IProgNo, IPreCalcOutput, IstormOutput
            INTEGER NHEADER, STAT, CREME 96 OPEN
            DATA NHEADER/3/
            CHARACTER*9 CREATION DATE
            CHARACTER*8 CREATION TIME
            CHARACTER*5 FEXT(10)
            DATA FEXT/'.GT1','.GT2','.GT3','.GT4','.GT5','.GT6','.GT7',
                     '.GT8','.GT9','.GTX'/
         &
   C-----
            CALL GET CREME96 VERSION (ICREME96 vno)
            IPreCalcOutput=0
           IF (PreCalcGTFs) THEN
             StormyPreCalc=.FALSE. !Local variable for header output file.
              IPreCalcOutput=1 !Local variable for header output file.
              IF ( (IpreCalc .EQ. 1) .OR. (IpreCalc .EQ. 3) )
                    StormyPreCalc=.TRUE.
           ENDIF
           IStormOutput=0
           IF (StormyPreCalc .OR. Stormy) IStormOutput=1
<sup>}</sup> c
           ILbins = 0 & ILbins = 1 from input routine are treated
TU c
           as ILbins = 1 for output, since they are stored in the
           same location in the array.
           IF (ILbins .EQ. 1 .AND. XLBOUNDS(1) .EQ. 0.0) THEN
                OPEN(UNIT=16,STATUS='NEW',FILE='USER:'//GtransFile)
               stat = creme96 open(gtransfile,'user',16,'new')
               CALL DATE (CREATION DATE)
               CALL TIME (CREATION TIME)
               WRITE(16,403) NHEADER, GTRANSFILE(1:70),
        #
                             ICREME96vno, IProgno
               WRITE(16,992) ICREME96vno, CREATION DATE, CREATION TIME
               WRITE(16,404) OrbIncl, Apogee, Perigee, AscNodeLong,
        #
                             AscNodeDisp, PerigDisp
               WRITE(16,405) IStormoutput, IPrecalcOutput, Year,
                             XLbounds (ILbins), XLbounds (Ilbins+1)
               DO I=1, Nrigs
                 WRITE (16,410) RigBins (I), TransFunc (I, ILbins)
               ENDDO
               CLOSE (16)
```

ELSE

```
TEMPFILE=Gtransfile(1:index(gtransfile,' ')-1)//FEXT(L)
                 OPEN(UNIT=16,STATUS='NEW',FILE='USER:'//GtransFile//FEXT(L))
     С
                stat = creme96_open(tempfile,'user',16,'new')
                CALL DATE (CREATION DATE)
                CALL TIME (CREATION TIME)
                WRITE(16,403) NHEADER, TEMPFILE(1:70),
         #
                              ICREME96vno, IProgno
                WRITE(16,992) ICREME96vno, CREATION_DATE, CREATION_TIME
                WRITE(16,404) OrbIncl, Apogee, Perigee, AscNodeLong,
         #
                              AscNodeDisp, PerigDisp
                IF (L .LT. NLvals) THEN
                    WRITE(16,405) IStormoutput, IPrecalcOutput, Year,
                              XLbounds(L), XLbounds(L+1)
                ELSE
                    WRITE(16,405) IStormoutput, IPrecalcOutput, Year,
                              XLbounds (L), XLinfinite
                ENDIF
                DO I=1, Nrigs
                  WRITE(16,410)RigBins(I),TransFunc(I,L)
                ENDDO
CLOSE (16)
              ENDDO
            ENDIF
W
            RETURN
ļ.
     403
            FORMAT(I3,1x,A70,I4,1x,I1)
33
     404
            FORMAT(1x,'%Incl = ',F7.3,' deg Apo = ',E10.4,
į.
                      ' Peri = ',E10.4,' km',1x,3(F6.2,1x))
        #
405
            FORMAT(1x,'%ISTORM =', I2,' IPRECALC =', I2,
' Grid Epoch = ',F6.1,' L Bin: ',2(E10.4,1X))
LI I
     991
            FORMAT(1x, A79)
     992
           FORMAT(1x,'%Created by CREME96:GTRANS_DRIVER Version', I4,
                      ' on ',A9,' at ',A8)
           FORMAT (5X, F6.3, 5X, E10.4)
     410
           END
                 !OutputTransFunc routine
   C-----
           SUBROUTINE GetPreCalcGTF(IPreCalc,RigBins,TransFunc)
    C
           IMPLICIT NONE
           INTEGER I, IPreCalc, Nrigs, NLvals
           PARAMETER (Nrigs=1001, NLvals=10)
           LOGICAL PreCalcInit
   C
           contain output rigidity vs. transmission function
           REAL RigBins(Nrigs), TransFunc(Nrigs, NLvals)
```

DO L=1, ILbins

```
REAL RIGPC (Nrigs)
        REAL PCGTF1 (Nrigs), PCGTF2 (Nrigs), PCGTF3 (Nrigs), PCGTF4 (Nrigs)
        COMMON/PreCalcCMN/PCGTF1, PCGTF2, PCGTF3, PCGTF4
        DATA PreCalcInit/.FALSE./
С
        Initialize pre-calculated GTFs
        IF (.NOT. PreCalcInit) THEN
          CALL InitPreCalcs (RigBins)
          PreCalcInit=.TRUE.
        ENDIF
C
        Set these each time, providing capability to change
C
        if input is so structured
        IF (IpreCalc .EQ. 0) THEN
         DO I=1, Nrigs
          TransFunc(I,1)=PCGTF1(I)
         ENDDO
       ENDIF
       IF (IpreCalc .EQ. 1) THEN
         DO I=1, Nrigs
          TransFunc(I,1)=PCGTF2(I)
         ENDDO
       ENDIF
       IF (IpreCalc .EQ. 2) THEN
         DO I=1, Nrigs
          TransFunc(I,1)=PCGTF3(I)
         ENDDO
       ENDIF
       IF (IpreCalc .EQ. 3) THEN
         DO I=1, Nrigs
          TransFunc(I,1)=PCGTF4(I)
         ENDDO
       ENDIF
       RETURN
       END
C-----
       SUBROUTINE CALCULATE TRANS FUNC (ISTEP, MAT, CF, NperLbin, T)
С
С
       Modified to allow the transmission function to be nonzero for
С
       C=0 bin (0.0-0.2 GV at present) , 1-29-96, PRB.
C
       Modified MAT to be real, in order to handle geometric shadowing
C
       for non circular orbits in a consistent manner
C
       Removed check on NstepSum being equal to ISTEP, since the minimum
       and maximum L-values of the specified bins do not have to
С
```

```
IMPLICIT NONE
                                         INTEGER ISTEP, J, Nrigs, NLvals, NstepSum, L
                                        PARAMETER (Nrigs=1001, NLvals=10)
                                        INTEGER NperLbin(NLvals)
                                        REAL MAT (Nrigs, NLvals)
                                        REAL T(Nrigs, NLvals), CF(Nrigs), CMAT(NLvals)
              C
                                       NstepSum=NperLbin(1)
                                       DO L=1, NL vals
                                              CMAT(L)=0.
                                              IF (L .GE. 2) NstepSum=Nstepsum+NperLbin(L)
                                       ENDDO
             C
                                       DO L=1, NL vals
                                              IF (NperLbin(L) .GT. 0) THEN
The line was from the line and 
                                                   DO J=1, Nrigs
            С
            С
                                             Convert the histogram to transmission.
                                                          CMAT(L) =MAT(J, L) / FLOAT(NperLbin(L)) + CMAT(L)
                                                          IF (L .EQ. 1) CF(J) = FLOAT(J-1) *0.02
                                                          T(J,L) = CMAT(L)
ini.
S
                                                    ENDDO
il de la cia
ENDIF
Ш
                                      ENDDO
ļ.
           C
20.9
                                      RETURN
                                      END
                                      SUBROUTINE NYMMIK(ReffGrid, TimeLocal, DeltaNymmik)
            C
                                      PURPOSE: To calculate Nymmik parameterization of Cutoff for
            C
                                      Rigidities below 1 GV, given the IGRF model. This attempts
            С
                                      to account for the quiet external magnetospheric fields for
            С
                                     high latitudes. The 1980 Shea & Smart 5 deg. by 5 deg. grid is
            C
                                      expected for the IGRF model. Uses standard local time as
                                     geomagnetic local time
                                      IMPLICIT NONE
                                     REAL ReffGrid, ReffNymmik, DeltaNymmik
```

REAL TimeLocal, HoursLT

REAL PI

include all L-values in the orbit. 11-21-96, PRB.

С

С

```
PI=ACOS(-1.0)
                          HoursLT= TimeLocal/3600.
         C
                          Reset correction to zero in case grid is zero
                          DeltaNymmik = 0.
         C
                            ReffNymmik=ReffGrid !handled in Geomag.
                          IF (ReffGrid .LE. 1.0 .AND. ReffGrid .GT. 0.) THEN
                              DeltaNymmik=1.42*(0.67/ReffGrid)**(1.1+
                                                           1.62*COS(2*PI/24*(HoursLT+0.6)))
                   >
                              DeltaNymmik=DeltaNymmik/EXP(1.72*(ReffGrid/0.67)**2*
                                                       (1+0.66*SIN(2*PI/12*(HoursLT+2.0))))
        C
                              Actual correction performed in SUBROUTINE Geomag
        C
        C
                                ReffNymmik=ReffGrid/(1+DeltaNymmik)
                         ENDIF
                         RETURN
The first time that the same t
                         END
                                         ------
                         SUBROUTINE Orbit (n, time, zlon, zlat, radius, alta, altp, a1, a2, a3, xi)
                         THIS SUBROUTINE USED TO ACCEPT INPUT CONCERNING SATELLITE
                         ORBITS AND CALCULATES THEIR GEOGRAPHICAL LOCATION.
EE
       C
                         N=0: disabled here, since the driver program now establishes
С
                                        all initial values.
C
                        N=1:
                                        Initialize data on orbit (complete mode).
                         N=2:
                                        CALCULATE ORBIT AS A FUNCTION OF TIME.
i
i
      C
                         Clearly data must be input before computations.
Q
       C
                         On data input, time returns the orbital period.
       C
                         During orbit calculations, time is an input variable.
       C
                         The following data should be passed into this Subroutine.
                         Re is now contained in a DATA statement. Only E (the eccentricity)
       C
                         is calculated in this version.
       C
       C
       C
                         ALTA=Orbital altitude at apogee (kilometers).
       C
                         ALTP=Orbital altitude at perigee (kilometers).
       C
                         RE=Radius of Earth (kilometers).
       C
                        E=Orbit eccentricity.
       С
                         al=Orbital inclination (degrees).
       C
                        A2=Initial longitude of ascending node (degrees).
       C
                        A3=Initial displacement from ascending node (degrees).
       C
                        XI=Displacement of perigee from ascending node (degrees).
       C
                         IMPLICIT REAL (A-H,O-Z)
                         INTEGER N
                        REAL pi,alta,altp,Re,e,a1,a2,a3,xi,w1,rmaj,w2,fact
                        REAL tho, pho, psi, xio
```

REAL time, zlon, zlat, radius

```
INTEGER I1014STEPS
        REAL DELMAX
C
        DATA Re/6371.2/
        pi=4.0*ATAN(1.0) !use ARCTAN to calculate PI
        IF (N.EQ.2) GO TO 1000
        E= (ALTA-ALTP) / (ALTA+ALTP+2.*RE)
        IF (E.LT..00001) E=0.
С
С
        W1=ANGULAR VELOCITY OF EARTH (RADIANS/SEC).
C
        RMAJ=SEMI-MAJOR AXIS (KILOMETERS).
C
        W2=MEAN ORBITAL ANGULAR VELOCITY (RADIANS/SECOND).
C
        TIME=ORBITAL PERIOD (SECONDS).
С
        FACT=A USEFUL FACTOR.
C
        W1=7.27E-5
        RMAJ = (ALTP + RE) / (1.-E)
        W2=1.24E-3*(RE/RMAJ)**1.5
        TIME=2.*PI/W2
        FACT=SQRT((1.+E)/(1.-E))
С
C
        DEFINE MORE USEFUL ANGLES.
        THO=PI*A1/180.
        PHO=PI*(A2-90.)/180.
        PSI=PI*A3/180.
        XIO=PI*XI/180.
        RETURN
                  !Finished initializing orbital variables
1000
        CONTINUE
С
C
        COMPUTE SATELLITE POSITION.
С
С
        QM=MEAN ANOMALY.
C
        IF (E.NE.O.) GOTO 1009
          QM=W2*TIME+PSI
        GOTO 1010
 1009
        CONTINUE
          YSS=(PSI-XIO)/2.
          QE0=2.*ATAN2(SIN(YSS), FACT*COS(YSS))
          QM0=QE0-E*SIN(QE0)
          QM=W2*TIME-QM0
 1010
        CONTINUE
C
        QE=ECCENTRIC ANOMALY.
C
        QE=QM
       DEL=1.
```

I1014STEPS=0 DELMAX=.0001

CONTINUE

1014

```
QTEMP=QE
              QE=QM+E*SIN(QE)
              DEL=QE-QTEMP
            IF (I1014STEPS .GE. 50) DELMAX=0.002
            IF (I1014STEPS .GE. 100) DELMAX=0.005
            IF (I1014STEPS .GE. 200) DELMAX=0.01
            IF (I1014STEPS .GE. 500) DELMAX=0.02
            IF (I1014STEPS .GE. 1000) DELMAX=0.05
            IF (ABS(DEL).GT.DELMAX .AND. I1014STEPS .LT. 2000) GOTO 1014
    C
    C
            QT=TRUE ANOMALY.
    C
            IF (E.NE.O.) GOTO 1019
              QT=QE
            GOTO 1020
            CONTINUE
     1019
              OECYC=INT(QE/2./PI)
              QERED=QE-2.*PI*QECYC
              AA1=FACT*SIN(QERED/2.)
              AA2=COS (QERED/2.)
              QTRED=2.*ATAN2(AA1,AA2)
              IF (OTRED.LT.0.) QTRED=QTRED+2.*PI
              QT=QTRED+2.*PI*QECYC+XIO
   C
            NOTE: ANOMALY COMPUTATIONS ARE DONE FROM PERIGEE
            WHILE ORBIT COMPUTATIONS BELOW ARE DONE FROM THE
   C
   C
            ASCENDING NODE. THE FACTOR OF XIO CORRECTS THIS.
   C
    1020
            CONTINUE
   С
   С
            ZLAT=LATITUDE.
FL
            R1=SIN(THO)*SIN(QT)
            THP=ACOS(R1)
ini.
            ZLAT=90.-180.*THP/PI
   С
   С
            ZLON=LONGITUDE.
    С
            RP=.5*(PI/2.+THO)
            RM=.5*(PI/2.-THO)
            RF = .5*(PI/2.-QT)
            IF (SIN(RF).NE.O.) GOTO 1029
              PHP=PI+PHO-W1*TIME
            GOTO 1030
     1029
            CONTINUE
              S1=SIN(RM) *COS(RF)/SIN(RP)/SIN(RF)
              S2=COS (RM) *COS (RF) /COS (RP) /SIN (RF)
              SUM=ATAN(S1)+ATAN(S2)
              PHP=PHO-W1*TIME+SUM
     1030
            CONTINUE
            IF (PHP.GE.O.) GOTO 1034
              PHP=PHP+2.*PI
            GOTO 1030
     1034
            CONTINUE
            IF (PHP.LT.2.*PI) GOTO 1035
              PHP=PHP-2.*PI
```

Ŧ

GOTO 1034

I1014STEPS=I1014STEPS+1

1035 CONTINUE
ZLON=180.*PHP/PI

C
C RADIUS=ALTITUDE (KILOMETERS).

C
RADIUS=RMAJ*(1.-E*COS(QE))-RE
RETURN !Actual orbital step computations
END

#

```
PROGRAM GtransDriver
С
        This program calculates the transmission functions as proposed in
С
С
        in the first year of the SEE work. It is intended as a driver
C
        program for subroutines which will be used in the overall program.
C
C
        At present, it does not allow for repeated calls. The overall
C
        structure should allow repeated calls, but would need to be tested.
C
        IMPLICIT NONE
        INTEGER Nrigs, NLvals
        PARAMETER (Nrigs=1001, NLvals=10)
С
        Rigidities expected in 0.02 GV steps & common for all L-bins
        REAL Transfunc(Nrigs, NLvals), RigBins(Nrigs), XLbounds(NLvals)
        Parameters that are input or initialized in GTFDriverInput
С
C
        These need to be passed to GEOMAG3.
        REAL OrbIncl, Apogee, Perigee, AscNodeLong, AscNodeDisp, PerigDisp
        REAL Zenith, Azimuth, UTtimeInit, Year
        INTEGER IpreCalc, ILbins
        LOGICAL Shadow, Stormy, PreCalcGTFs
        CHARACTER*80 GtransFile
        INTEGER Iprogno
        DATA IprogNo/2/
        CALL GTFDriverInput (OrbIncl, Apogee, Perigee, AscNodeLong,
     #
               AscNodeDisp, PerigDisp, Zenith, Azimuth, UTtimeInit, Stormy,
     #
                Shadow, PreCalcGTFs, IPreCalc, GtransFile, Year, XLbounds,
     #
                ILbins)
        CALL Geomag96 (Orbincl, Apogee, Perigee, AscNodeLong, AscNodeDisp,
     #
                 PerigDisp, Zenith, Azimuth, UTtimeInit, Stormy, Shadow,
     #
                 PreCalcGTFs, IPreCalc, RigBins, TransFunc, Year, XLbounds,
     #
                 ILbins)
C
        For adding header information to output GTF file. Added July 1996.
        CALL OutputTransFcn(RigBins, TransFunc, GtransFile, OrbIncl, Apoque,
                Perigee, AscNodeLong, AscNodeDisp, PerigDisp, Zenith, Azimuth,
     #
     #
               UTtimeInit, Stormy, Shadow, PreCalcGTFs, IPreCalc, Year,
     #
               XLbounds, ILbins, IprogNo)
        STOP
        END
        SUBROUTINE GTFDriverInput(OrbIncl, Apogee, Perigee, AscNodeLong,
     #
                       AscNodeDisp, PerigDisp, Zenith, Azimuth, UTtimeInit,
```

Stormy, Shadow, PreCalcGTFs, IPreCalc, GtransFile,

IMPLICIT NONE

REAL OrbIncl, Apogee, Perigee, AscNodeLong, AscNodeDisp, PerigDisp REAL Zenith, Azimuth, UTtimeInit

Note that the eccentricity is calculated here to decide if
need to read PerigDisp. The eccentricity is also recalculated
in the initialization CALL ORBIT(1,...) case found in SUBROUTINE GEOMAG
This makes the input driver independent of the actual computational
routines, so that it will be easier to modify and interface with other
space environment routines.

REAL E,Re !eccentricity and radius of Earth
PARAMETER (Re=6371.2)
LOGICAL Shadow,Stormy,PreCalcGTFs
REAL ApPerSwitch

INTEGER Istorm,Ishadow,IPreCalc
INTEGER Itype,IGTFtype !look-out directions & pre/non-pre calc. GTFs
INTEGER IERR,IACCEPT
DATA IERR/0/

CHARACTER*80 GtransFile

INTEGER NLvals,I,L,ILbinMax,ILbinsum
PARAMETER (NLvals=10)

REAL XLbounds(NLvals), XLinfinite, Year PARAMETER (XLinfinite=1.0E+06)

REAL XLdummy

C-----

WRITE(6,1000)
WRITE(6,1001)

UTtimeInit=0.0 !start at 0 UT by default

C Present averaging algorithms assume that zenith & azimuth correspond to vertical incidence.

Zenith=0.0 Azimuth=0.0

C initialize boundaries L-value bins

XLbounds(1)=0.0
Year=1980.0 !needed for L-value calculations

DO L=2,NLvals
XLbounds(L)=XLinfinite
ENDDO

C Check if user wants to use a pre-calculated GTF. If so, read specified option and return.

9390 CONTINUE

```
CALL RETRY INPUT (IERR)
         WRITE(*,390)
         READ (*, *, ERR=9390, IOSTAT=IERR) IGTFtype
         PreCalcGTFs=.FALSE. !initialize not to use pre-calculated GTFs
C
        Note that pre-calculated GTFs already include Earth shadow,
C
         since solid Earth is included in the trajectory-tracing calculations.
C
         In those cases, Shadow must be set to .FALSE. regardless of user
C
         input.
        IF (IGTFtype .NE. 0) THEN
 9391
           CONTINUE
           CALL RETRY_INPUT(IERR)
           WRITE(*,391)
           READ(*,*,ERR=9391,IOSTAT=IERR)IpreCalc
           PreCalcGTFs=.TRUE.
           Shadow=.FALSE.
C
          Use quiet-time, 51.6 degrees as the default case
           IF (IpreCalc .LT. 0 .OR. IpreCalc .GT. 3) IpreCalc=0
 9427
          CONTINUE
          CALL RETRY INPUT (IERR)
          WRITE(*,427)
          READ(*,428,ERR=9427,IOSTAT=IERR)GtransFile
          CALL CHECK_OUTPUT_FILE(Gtransfile,IACCEPT)
          IF (IACCEPT.NE.0) GOTO 9427
С
          For use in SUBROUTINE GTFHeaderOutput. Added July 1996.
          IF (IpreCalc .EQ. 0 .OR. IpreCalc .EQ. 1) THEN
            OrbIncl=51.6
            Apogee=450.0
            Perigee=450.0
          ELSEIF (IpreCalc .EQ. 2 .OR. IpreCalc .EQ. 3) THEN
            OrbIncl=28.5
            Apogee=450.0
            Perigee=450.0
          ENDIF
C
          The pre-calculated GTFs are not presently divided into L-bins
          ILbinsum=1
          RETURN
        ENDIF
C
        Hardwire shadow to be TRUE
        Shadow=.TRUE.
C
        Choose from the two original CREME options for the state of the
C
C
        magnetosphere (quiet or stormy). Note the stormy option applies on
С
        top of the Nymmik correction for mid to high-latitudes.
        CONTINUE
9412
        CALL RETRY INPUT (IERR)
        WRITE (*,412)
        READ (*,*,ERR=9412,IOSTAT=IERR) Istorm
```

Stormy=.FALSE.

```
IF (Istorm .EQ. 1) Stormy=.TRUE.
 C
         What is the altitude at apogee?
C
  9420
         CONTINUE
         CALL RETRY INPUT (IERR)
         WRITE(*,420)
         READ (*,*,ERR=9420,IOSTAT=IERR) Apogee
С
C
         WHAT IS THE ALTITUDE AT PERIGEE?
C
 9400
         CONTINUE
         CALL RETRY INPUT (IERR)
        WRITE (*, 400)
        READ (*,*,ERR=9400,IOSTAT=IERR) Perigee
С
        allow the user to specify apogee and perigee in either order
С
        instead of performing unintended calculation which sets eccentricity
С
        to zero and using Perigee variable (actual apogee) to produce
C
        a circular orbital altitude in ORBIT routine.
        IF (Perigee .GT. Apogee) THEN
           ApPerSwitch=Apogee
           Apogee=Perigee
           Perigee=ApPerSwitch
           WRITE(*,430)
        ENDIF
        E= (Apogee-Perigee) / (Apogee+Perigee+2.*Re)
        IF (E.LT..00001) E=0.
C
С
        WHAT IS THE ORBITAL INCLINATION?
C
 9405
        CONTINUE
        CALL RETRY INPUT (IERR)
        WRITE(*,405)
        READ (*,*,ERR=9405,IOSTAT=IERR)
                                          OrbIncl
C
        Have Removed "FAST" option, i.e. must enter Ascending Node information
C
С
        Retain these initializations in case want to hardwire ascending
        node information at future time.
        AscNodeLong=0.
        AscNodeDisp=0.
        PerigDisp=0.
C
        WHAT IS THE INITIAL LONGITUDE OF THE ASCENDING NODE?
C
        WRITE(*,409)
 9410
        CONTINUE
        CALL RETRY INPUT (IERR)
        WRITE(*,410)
        READ (*,*,ERR=9410,IOSTAT=IERR) AscNodeLong
C
C
        WHAT IS THE INITIAL DISPLACEMENT FROM THE ASCENDING NODE?
C
```

```
CALL RETRY INPUT (IERR)
            WRITE(*,415)
            READ (*, *, ERR=9415, IOSTAT=IERR) AscNodeDisp
            IF (E.NE.O.) THEN
                                    !Only read in XI if eccentricity is nonzero
    C
   С
                What is the displacement of the perigee from the ascending node?
   C
    9425
             CONTINUE
             CALL RETRY INPUT (IERR)
             WRITE (*, 425)
             READ (*, *, ERR=9425, IOSTAT=IERR) PerigDisp
           ENDIF
           IF ( (AscNodeLong .NE. 0.0) .OR. (AscNodeDisp .NE. 0.0) .OR.
                 (PerigDisp .NE. 0.0) ) WRITE(*,426)
            Itype = 1
                        !hardwire vertical incidence, applied with shadow.
    9450
           CONTINUE
           CALL RETRY_INPUT(IERR)
           WRITE(*,450)
           READ (*,*,ERR=9450,IOSTAT=IERR)
                                             ILbinMax
           IF (ILbinMax .LT. 0) ILbinMax=0
           IF (ILbinMax .GT. NLvals) THEN
WRITE(*,456)
1
               ILbinMax = NLvals
           ENDIF
Ξ
           IF (ILbinMax .GT. 0) THEN
WRITE(*,451) ILbinMax
9451
               CONTINUE
W
               CALL RETRY INPUT (IERR)
į.
               READ (*, *, ERR=9451, IOSTAT=IERR) (XLbounds(L), L=1, IlbinMax)
                IF (ILbinMax .EQ. 1 .AND. XLBOUNDS(1) .EQ. 0.0) WRITE(*,458)
           ENDIF
           IF (XLbounds(1) .LT. 0.0) XLbounds(1)=0.0
   C
           Start DO loop at 1, so that ILbinMax=2 will be properly handled
           This SUBROUTINE insists the L-values are in increasing order.
   С
   C
           If this is not the case, all subsequent L-value bins will be
   C
           ignored.
           DO L=1, ILbinMax
             IF (XLbounds(L) .LT. XLbounds(1)) THEN
                 WRITE(*,452) XLbounds(L), XLinfinite
                 XLbounds (L) =XLinfinite
             ENDIF
             IF (L .GE. 2) THEN
               IF (XLbounds(L) .LE. XLbounds(L-1)) THEN
                 WRITE(*,452) XLbounds(L), XLinfinite
                 XLbounds (L) =XLinfinite
               ENDIF
             ENDIF
```

9415

CONTINUE

ENDDO

```
ILbinsum=1
       DO L=1, ILbinMax
         IF ( (L .GE. 2) .AND. (XLbounds(L) .LT. XLinfinite) )
               ILbinsum=ILbinsum+1
       ENDDO
       IF (ILbinMax .NE. ILbinsum .AND. ILbinMax .NE. 0) THEN
         WRITE (*, 453) ILbinMax, ILbinsum
         ILbinMax=ILbinsum
       ENDIF
       IF ( ILbinMax .GT. 1 .OR. (ILbinMax .EQ. 1 .AND.
           XLBOUNDS(1) .GT. 0.0) ) THEN
9454
         CONTINUE
         CALL RETRY INPUT (IERR)
         WRITE(*,454)
         READ (*,*,ERR=9454,IOSTAT=IERR) Year
       ENDIF
9428
       CONTINUE
       CALL RETRY INPUT (IERR)
       IF (ILbinMax .EQ. 0 .OR. (ILbinMax .EQ. 1 .AND.
           XLBOUNDS(1) .EQ. 0.0) ) THEN
    &
         WRITE(*,427)
         READ (*, 428, ERR=9428, IOSTAT=IERR) GtransFile
         CALL CHECK OUTPUT FILE (Gtransfile, IACCEPT)
         IF (IACCEPT.NE.0) GOTO 9428
       ELSE
         WRITE(*,455)ILbinMax
         READ(*,428,ERR=9428,IOSTAT=IERR)GtransFile
         CALL CHECK OUTPUT FILE (Gtransfile, IACCEPT)
         IF (IACCEPT.NE.0) GOTO 9428
        DO I=1, LEN (GtransFile)
           IF (GtransFile(I:I) .EQ. '.') THEN
               GtransFile=GtransFile(1:I-1)
           ENDIF
         ENDDO
       ENDIF
       RETURN
1000 FORMAT(1X,'GEOMAG96 Geomagnetic Transmission Function Model',/)
1001 FORMAT(' This program will calculate the omnidirectional',
    &
             ' geomagnetic transmission',
           /,' function (GTF) to a',
    &
              ' spacecraft orbiting inside the magnetosphere. The',
    &
           /,' calculated GTF is used by the CREME96',
             ' particle environment model.',
    &//,' NOTE: Before running this or any other CREME96 programs',
    & ' please define three ',
    & /,' logicals: ',/,
    & /,4x,' CREME96
                           as the directory where CREME96 source',
             ' & executables reside.',
    & /,4x,' CR96TABLES
                           as the directory in which CREME96 data',
             ' tables reside.',
    & /,4x,' USER
                           as the directory in which output files',
             ' should be written.',
```

```
& //,' Now begin specification of the GTF calculation: ',/)
C
                                2
                                          3
             12345678901234567890123456789012345678901234567890
С
 390
        FORMAT(1X,'Enter 1 in order to use a pre-calculated GTF for a',
                  ' typical space shuttle or',
    & /,3x,' space station orbit, ie., 28.5 deg or 51.6 deg & 450 km.'
     & //,1X,'Enter 0 to specify an arbitrary orbit: ',
     & //,3X,'[The pre-calculated GTFs are recommended if appropriate,',
            ' since these use a'
    & /,3x,' better magnetic field model than used in the arbitrary',
            ' orbit option.]')
391
       FORMAT(1X,'Enter 0 for Space Station (51.6 deg., 450 km)',
             ' orbit (ISSA), quiet magnetosphere',
    & /,7X,'1 for ISSA, stormy magnetosphere',
    & /,7X,'2 for 28.5 deg. (450 km), quiet magnetosphere',
    & /,7X,'3 for 28.5 deg. (450 km), stormy magnetosphere: ',
    & //,3X,'[(a) For solar-quiet periods, the quiet magnetosphere',
              ' is typical.',
        /,3X,' (b) For solar energetic particles, the stormy',
    &
    &
              'magnetosphere should also'
    & /,8X,'be considered.]',
    & //,3X,'NOTE: the Worst Day in 22 years (see the CREME96',
            ' environment model)'
    & /,3X,'included a stormy magnetosphere,',
            ' and thus a stormy option must be considered',
    & /,3X,'with this Worst Day option.')
412
      FORMAT(/,1X,'Enter the magnetospheric field condition: 0',
    # 'for quiet; 1 for stormy: ',
    & //,3X,'[(a) For solar-quiet periods, the quiet magnetosphere',
             ' is typical.',
    & /,3X,' (b) For solar energetic particles, the stormy',
             'magnetosphere should also'
    &
    & /,8X,'be considered.]',
    & //,3X,'NOTE: the Worst Day in 22 years (see the CREME96',
            ' environment model)'
    &
    & /,3X,'included a stormy magnetosphere.',
          ' For many orbits, the generic stormy GTF'
    &
    & /,3x,'calculated here',
            ' can be substantially smaller than the actual GTF.')
420
       FORMAT(/,1X,'Enter altitude at apogee (kilometers): ')
400
       FORMAT(/,1X,'Enter altitude at perigee (kilometers): ')
430
       FORMAT(/,1X,'Input apogee < perigee, have been interchanged.')
405
       FORMAT(/,1X,'Enter orbital inclination (degrees): ')
       FORMAT(/,1X,'The remaining input parameters are most relevant',
409
       ' to situations in which the',
    ₽.
    & /,1X,'actual orbital path is known',
            ' or in which mission critical operations are',
    & /,1x,'planned.',
    & //,3X,' [Recommended values are 0.0, unless you wish to examine',
    & /,3X,'a very specific orbital segment.]')
410
       FORMAT(/,1X,'Enter initial longitude of ascending node',
                1X, '[Recommended = 0.0 (degrees)]:')
415
       FORMAT(/,1X,'Enter initial displacement from ascending',
```

```
' node', 1X, '[Recommended = 0.0 (degrees)]:')
     FORMAT(/,1X,'Enter displacement of perigee from',
    1 'ascending node',1X,'[Recommended = 0.0 (degrees)]:')
       FORMAT(/,1X,'Note: for studies sensitive to a specific',
    & 'orbital segment, you should be',
    & /,1X,'aware that the GTF',
    & ' calculations are averaged over 7 days at present. This',
    & /,1X, 'parameter can be easily reset by modifying',
    & 'the GEOMAG96 subroutine, but is ',
    & /,1x,'not provided as a general-use input parameter.')
427
       FORMAT(/,1X,'Enter name of output GTF file:',
             /,' [Recommended: something.GTF]')
      FORMAT (A80)
428
450
      FORMAT(/,1X,'Enter the number of desired GTF L-value bins ',
       '(1 - 10):',
    & /,3X,'[Recommended default = 0, i.e.',
            'one GTF for the entire orbit.]')
451 FORMAT (/, 1X,
           'Enter the lower limits of the ', I2,' L-value bins: ',
    & /,3X,'[A typical scenario could be to request 4 bins as the',
         ' as the previous entry.',
   & /,3X,'Then, entries of 0.0, 2.0, 4.0, and 6.0',
           ' would subdivide the orbit into',
   & /,3x,'sections with L < 2, L = 2-4, L = 4-6, and L > 6.]',
   & //,1X,'NOTE: The L-value is a magnetic coordinate roughly',
          ' corresponding to the',
   & /,1X,'distance in Earth Radii to the',
   & /,1x,'For example, a geosynchronous orbit is roughly L = 6.6,',
          ' the geographic equator',
   &
   & /,lx,'is about L = 1, and the heart of the',
   &
          ' South Atlantic Anomaly (SAA) is roughly at',
   \& /,1x,'L = 1.2 - 2.',
   & ' Calculated L-values slightly less than 1 do occur; using',
   & /,1X,'a lower limit of L = 0 will account for these.')
452 FORMAT(1X,'The L-values MUST be entered in increasing order',
   & /,1X,'the L-value of ',F10.2,' has been reset to ',F10.2)
      FORMAT(1X,'The number of L-values bins has been reset',
453
   & /,1X,' from ',I2,' to ',I2)
454
      FORMAT(/,1X,
           'Enter the decimal year for the field model in the ',
   &
           'L-value calculations:',
   & /,1X,' [Since the present IGRF grid calculations were performed',
           ' for 1980.0, that date',
   & /,1X,' is presently recommended for consistency.]')
    FORMAT(/,1X,'Enter root name of output GTF files:',
   & /,1X,'[NOTE: There will be ',I2,' output files, and',
   & 'the files for the different L-value'
   & /,1x,' bins will',
           ' be called something.GT# (# = 1,2,...,9,X)]')
```

```
FORMAT(1X,'Only 10 L-values are allowed.')

FORMAT(1X,'Calculation reset to whole orbit option, since',

X,'choosing 1 L bin',

/,1X,' with a minimum L-value equal to 0 is equivalent to',

1X,'the entire orbit.')

END !GTFDriverInput routine
```

```
the state of the s
```

```
SUBROUTINE HEAVY_ION_UPSETS (LET_FILE, XM, YM, ZM, FUNNELM,
      &
                                      IPARAM, PARAMS,
                                      XSECT_FILE, NBITS, IENTER,
      &
      &
                                      SEU RATE, DAY RATE, PERSECOND, PERDAY)
C
C
         Subroutine for performing heavy-ion evaluation:
С
         Inputs: LET FILE = file containing integral LET spectrum
C
                                 (in ions/m2-s-sr) vs. LET (in MeV-cm2/g)
C
                  XM,YM,ZM = bit dimensions (in microns)
C
                  FUNNELM = funnel length (optional; default 0) in microns
С
                              = 1,2,4, indicating cross-section model
                  IPARAM
С
                                1 = Bendel 1-parameter
C
                                2 = Bendel 2-parameter
С
                                4 = Weibull
С
                                5 = Critical charge (pc)
С
                                0 = table
C
                                  = array containing cross-section parameters
                  PARAMS (4)
C
                  XSECT FILE = file containing cross-section table.
С
                  NBITS
                                  = no. bits in the device:
С
C
        Outputs: SEU RATE
                               in SEUs/bit/second
C
                   DAY RATE
                               in SEUs/bit/day
C
                   PERSECOND
                               in SEUs/device/second
С
                               in SEUs/device/day
                   PERDAY
С
C
        Written by:
                        Allan J. Tylka
C
                        Code 7654
C
                        Naval Research Laboratory
C
                        Washington, DC 20375-5352
С
                        tylka@crs2.nrl.navy.mil
С
С
C--
С
        IMPLICIT NONE
        INTEGER*4 NBINS, NPTS, IPARAM, K, IENTER, MPTS
        REAL*4 LET, LETMG, FLUX, PARAMS, XM, YM, ZM, FUNNELM, XSECT, QC
        REAL*4 NBITS, SEU RATE
        REAL*4 DAY RATE, PERSECOND, PERDAY
        CHARACTER*80 LET FILE, XSECT FILE
        PARAMETER (NBINS=5000)
        DIMENSION LET (NBINS), LETMG (NBINS), FLUX (NBINS), XSECT (NBINS)
        DIMENSION PARAMS (4)
C
        WRITE(6,9998)
 9998
        FORMAT(1x,' HI UPSET DRIVER calculation started.',
                 ' Please stand by.')
        SEU RATE=0.0
C
        On first entry, get integral LET spectrum:
        IF (IENTER.EQ.1) THEN
        CALL UNLOAD LET SPECTRUM(LET FILE, LET, FLUX, NPTS)
```

C

```
С
         expects it in MeV-cm2/mg.
         DO 100 K=1, NPTS
            LETMG(K) = LET(K) \star0.001
 100
         CONTINUE
         ENDIF
         IF (XM*YM.GT.0.0) THEN
         IF (IPARAM.NE.5) THEN
C
             For devices in which the cross-section has not reached its
C
C
             limiting value at effective LET = LET(NPTS), the cross-section
C
             table must be extended to higher effective LETs to MPTS:
             CALL EXTEND EFFECTIVE LET RANGE (NPTS, NBINS,
     æ
                                               XM, YM, ZM, FUNNELM, MPTS,
     &
                                               LET, LETMG, FLUX)
С
             Evaluate SEU cross-section at these effective LET values:
             CALL EVALUATE_SEU_CROSS_SECTION(LETMG, MPTS, IPARAM, PARAMS,
     &
                                              XSECT FILE, XSECT)
C
             Calculate SEU rate:
             CALL INTEGRATE_HEAVY_ION_UPSETS (MPTS, LET, FLUX, XSECT,
     &
                                              XM, YM, ZM, FUNNELM,
     &
                                               SEU_RATE)
        ELSEIF (IPARAM.EQ.5) THEN
            QC=PARAMS (1)
             CALL GET_UPSET(XM, YM, ZM, FUNNELM, QC, NPTS, LET, FLUX, SEU_RATE)
C
            Re-scale to allow for possibility of different limiting
C
             cross-section, rather than the customarily used XM*YM
            SEU_RATE=SEU_RATE*PARAMS(2)/XM/YM
        ENDIF
        ENDIF
        IF (SEU RATE.LT.O.) THEN
            WRITE(6,999) SEU RATE
 999
            FORMAT(1x,' ERROR in HEAVY_ION UPSETS: SEU RATE = ',E13.5)
            SEU RATE=0.0
        ENDIF
        CALL CALC_SEU_RATE(NBITS, SEU_RATE, DAY_RATE, PERSECOND, PERDAY)
        WRITE(6,9999)
        FORMAT(1x,' HI_UPSET_DRIVER calculation completed. ')
 9999
```

LET in spectrum file is in MeV-cm2/g; but cross-section evaluation

```
SUBROUTINE EXTEND EFFECTIVE LET_RANGE(NPTS, NBINS,
                                                 XM, YM, ZM, FUNNELM, MPTS,
     &
                                                 LET, LETMG, FLUX)
     &
        Based on device dimesions, extends the range of effective LET
C
        values from NPTS to MPTS
С
C
        IMPLICIT NONE
        INTEGER*4 NPTS, NBINS, MPTS
        REAL*4 XM, YM, ZM, FUNNELM
        REAL*4 LET, LETMG, FLUX
        DIMENSION LET(1), LETMG(1), FLUX(1)
        INTEGER*4 K, NLAST, NEXTRA
        REAL*4 FACTOR, DL
        MPTS=NPTS
        Locate last non-zero integral flux point:
C
        NLAST=0
        DO 100 K=1, NPTS
            IF (FLUX(K).LE.0.0 .and. NLAST.EQ.0) NLAST=K
 100
        CONTINUE
        FACTOR= (SQRT (XM*XM+YM*YM+ZM*ZM) +FUNNELM) /ZM
        DL=ALOG(LET(NPTS)/LET(1)) **(1./FLOAT(NPTS-1))
        NEXTRA=1.+ALOG(FACTOR)/ALOG(DL)
        MPTS=NEXTRA+NLAST
        IF (MPTS.GT.NBINS) MPTS=NBINS
        IF (MPTS.GT.NPTS) THEN
        DO 200 K=NPTS+1, MPTS
                LET (K) = LET (NPTS) *DL** (K-NPTS)
                LETMG(K) = LET(K) \star0.001
                FLUX(K) = 0.0
 200
        CONTINUE
        ENDIF
C
С
        Debug:
С
         TYPE *,' NPTS, LET(NPTS): ',NPTS,LET(NPTS)
         TYPE *,' NLAST, LET(NLAST): ',NLAST, LET(NLAST)
C
         TYPE *, ' FACTOR, NEXTRA, MPTS: ', FACTOR, NEXTRA, MPTS
С
         TYPE *,' LET (MPTS):
                                        ', LET (MPTS)
C
        RETURN
        END
```

```
PROGRAM HI UPSET_DRIVER
       IMPLICIT NONE
       REAL*4 XM, YM, ZM, FUNNELM, NBITS, PARAMS
       REAL*4 XMO, YMO, ZMO
       REAL*4 SEU RATE, DAY RATE, PERSECOND, PERDAY
       INTEGER*4 IPARAM, IREPEAT, IENTER
       DIMENSION PARAMS(4)
       CHARACTER*80 LET_FILE, XSECT_FILE, REPORT_FILE
       CHARACTER*40 DEVICE LABEL
       INTEGER*4 IERR
       DATA IERR/0/
       INTEGER*4 IENT
       DATA IENT/0/
C
       Modified 11/8/96: to extract XM, YM from cross-section data
C
                          if user supplies XM=YM=0.
C
C
       IENTER=1
       CONTINUE
 10
       CALL INITIALIZE_HI_UPSETS(LET_FILE,XM0,YM0,ZM0,FUNNELM,NBITS,
                                   IPARAM, PARAMS, XSECT_FILE, IENTER,
     &
                                   DEVICE LABEL, REPORT FILE)
     &
       CALL CHECK RPP DIMENSIONS (XMO, YMO, ZMO,
                                   IPARAM, PARAMS, XSECT_FILE,
     æ
                                   XM, YM, ZM)
     æ
       CALL HEAVY_ION_UPSETS(LET_FILE,XM,YM,ZM,FUNNELM,IPARAM,PARAMS,
                               XSECT FILE, NBITS, IENTER,
     &
                               SEU RATE, DAY RATE, PERSECOND, PERDAY)
     &
       CALL HI UPSET REPORT (LET FILE, XM, YM, ZM, FUNNELM, NBITS,
                              IPARAM, PARAMS, XSECT_FILE, IENTER,
     &
                              DEVICE LABEL, REPORT_FILE,
     &
                              SEU RATE, DAY_RATE, PERSECOND, PERDAY)
     &
 9100
            CONTINUE
            CALL RETRY INPUT (IERR)
            WRITE(6,9200)
            FORMAT(//,' Repeat SEU rate calculation with different',
 9200
                       ' device characteristics? (1=yes,0=no)')
            READ (*, *, ERR=9100, IOSTAT=IERR) IREPEAT
            IF (IREPEAT.EQ.1) THEN
                IENTER=IENTER+1
                GOTO 10
            ENDIF
       WRITE(6,9600)
 9600 FORMAT(1x,' Heavy Ion Upset Calculations finished.')
        STOP
        END
```

```
SUBROUTINE HI_UPSET_REPORT(LET_FILE, XM, YM, ZM, FUNNEL, NBITS,
                            IPARAM, PARAMS, XSECT FILE, IENTER,
    &
                            DEVICE LABEL, REPORT FILE,
    &
                            SEU RATE, DAY RATE, PERSECOND, PERDAY)
    æ
      IMPLICIT NONE
      REAL*4 XM, YM, ZM, FUNNEL, NBITS, PARAMS
      REAL*4 SEU RATE, DAY RATE, PERSECOND, PERDAY
      INTEGER*4 IPARAM, IENTER, OUTUNIT, VERSION_NUMBER, NHEADERO, K
      INTEGER*4 NHEADER, PROGRAM CODE, STAT, CREME96 OPEN
      DATA OUTUNIT/2/
      DIMENSION PARAMS (4)
      CHARACTER*80 LET FILE, XSECT_FILE, REPORT_FILE
      CHARACTER*40 DEVICE_LABEL
      CHARACTER*9 CREATION DATE
      CHARACTER*8 CREATION_TIME
      PROGRAM CODE=10
      IF (IENTER.EQ.1.and.REPORT_FILE.NE.'NULLFILE') THEN
        OPEN(UNIT=OUTUNIT,FILE='USER:'//REPORT FILE,STATUS='NEW')
       stat = creme96 open(report file, 'user', outunit, 'new')
       CALL DATE (CREATION_DATE)
       CALL TIME (CREATION TIME)
       CALL GET_CREME96_VERSION (VERSION_NUMBER)
       CALL CHECK HEADER LENGTH (LET FILE, NHEADERO)
       NHEADER=NHEADER0+2
       WRITE (OUTUNIT, 991) NHEADER, REPORT FILE (1:70),
                           VERSION NUMBER, PROGRAM_CODE
       FORMAT(I3,1x,A70,I4,I2)
991
       WRITE (OUTUNIT, 992) VERSION NUMBER, CREATION_DATE, CREATION_TIME
       FORMAT(1x,'%Created by CREME96:HI UPSET DRIVER Version', I4,
992
                ' on ',A9,' at ',A8)
       Now copy header information from input file:
       WRITE (OUTUNIT, 993) LET_FILE (1:40)
       FORMAT(1x, '%Input Integral LET Spectrum File: ', A40)
993
       CALL COPY HEADERS (LET FILE, NHEADERO, OUTUNIT)
      ENDIF
      IF (REPORT FILE.NE.'NULLFILE') THEN
       WRITE (OUTUNIT, 994) IENTER, DEVICE_LABEL
       FORMAT(/,1x,' REPORT NO. ',14,': ',2x,A40)
994
       WRITE (outunit, 995) XM, YM, ZM, FUNNEL
       FORMAT(1x,' RPP Dimensions: X = ', F10.5, ' Y = ', F10.5,
995
                    ' Z = ',F10.5,' microns.',
             /,1x,' Funnel length = ',F10.5,' microns.')
       IF (IPARAM.EQ.0) WRITE(outunit,980) IPARAM,XSECT_FILE(1:75)
       IF (IPARAM.EQ.1) WRITE(outunit, 981) IPARAM, PARAMS(1)
       IF (IPARAM.EQ.2) WRITE(outunit,982) IPARAM, PARAMS(1), PARAMS(2)
       IF (IPARAM.EQ.4) WRITE(outunit,984) IPARAM,(PARAMS(K),K=1,4)
       IF (IPARAM.EQ.5) WRITE(outunit, 985) IPARAM, PARAMS(1), PARAMS(2)
       WRITE (outunit, 996) NBITS
996
       FORMAT(1x,' Number of bits = ',E13.5)
       FORMAT(1x,' CROSS-SECTION INPUT ', I3,' FROM FILE: ',
980
             /,5x,A75)
       FORMAT(1x,' CROSS-SECTION INPUT', I3,
981
                 ' BENDEL 1-PARAMETER = ',E13.5)
```

```
FORMAT(1x, 'CROSS-SECTION INPUT', 13,
982
                ' BENDEL 2-PARAMETERS A,B = ',2E13.5)
   δe
      FORMAT(1x, ' CROSS-SECTION INPUT ', I3,
984
               ' WEIBULL FIT: ',
   &
           /,5x,'ONSET = ',F9.3,' MeV-cm2/milligram',
           /,5x,' WIDTH = ',F9.3,' MeV-cm2/milligram',
    &
           /,5x,' POWER = ',F9.3,' (dimensionless)',
    &
           /,5x,' PLATEAU = ',F9.3,' square microns/bit')
      FORMAT(1x,' CROSS-SECTION INPUT ',13,
985
           /,5x,' Critical charge = ',E13.5,' picocoloumbs',
    &
           /,5x,' Cross-Section = ',E13.5,' square microns/bit')
    &
      WRITE (outunit, 9200)
      WRITE(outunit, 9201) IENTER, SEU_RATE, DAY_RATE, PERSECOND, PERDAY
      FORMAT(2x,'Rates: SEUs/bit/second /bit/day',
9200
                 ' /device/second /device/day')
      FORMAT(2x,'*****', I4,2x,4(E14.5,2x))
9201
      ENDIF
      WRITE(6,9200)
      WRITE(6,9201) IENTER, SEU_RATE, DAY_RATE, PERSECOND, PERDAY
      RETURN
      END
```

```
IMPLICIT NONE
          Explicit variable lengths added AJT 12-12-96
    C
           INTEGER n, indx(n), M, NSTACK
    C
           REAL arr(n)
    C
           PARAMETER (M=7, NSTACK=50)
    С
           INTEGER i,indxt,ir,itemp,j,jstack,k,l,istack(NSTACK)
    С
    С
           REAL a
          Parameter NMAX added to define size of passed-in arrays.
    C
          INTEGER*4 n, NMAX, indx(1), M, NSTACK
          REAL*4 arr(1)
          PARAMETER (M=7, NSTACK=50)
          INTEGER*4 i,indxt,ir,itemp,j,jstack,k,l,istack(NSTACK)
          REAL*4 a
          IF (N.GT.NMAX) THEN
              WRITE(6,9999) N,NMAX
               FORMAT('3 99999 ABNORMAL TERMINATION: ',
     9999
                 /,lx,' Error in INDEXX: N,NMAX: ',2I12,
         &
                 /,1x,' STOP.')
               STOP
          endif
          do 11 j=1,n
            indx(j)=j
          continue
    11
          jstack=0
          1=1
           if (ir-1.lt.M) then
             do 13 j=1+1, ir
               indxt=indx(j)
               a=arr(indxt)
               do 12 i=j-1,1,-1
                 if(arr(indx(i)).le.a)goto 2
indx(i+1) = indx(i)
F.
               continue
    12
               i=0
l.A.
    2
               indx(i+1)=indxt
    13
             continue
             if (jstack.eq.0) return
             ir=istack(jstack)
             l=istack(jstack-1)
             jstack=jstack-2
           else
             k = (1 + ir)/2
             itemp=indx(k)
             indx(k) = indx(l+1)
             indx(1+1)=itemp
             if (arr(indx(l+1)).gt.arr(indx(ir)))then
               itemp=indx(l+1)
               indx(l+1)=indx(ir)
               indx(ir)=itemp
             endif
             if(arr(indx(l)).gt.arr(indx(ir)))then
               itemp=indx(1)
               indx(1)=indx(ir)
               indx(ir)=itemp
             endif
             if(arr(indx(l+1)).gt.arr(indx(l)))then
```

itemp=indx(1+1)
indx(1+1)=indx(1)

SUBROUTINE indexx(n,NMAX,arr,indx)

```
indx(1)=itemp
        endif
        i=1+1
        j=ir
        indxt=indx(1)
        a=arr(indxt)
3
        continue
          i=i+1
        if(arr(indx(i)).lt.a)goto 3
        continue
4
          j=j-1
        if(arr(indx(j)).gt.a)goto 4
        if(j.lt.i)goto 5
        itemp=indx(i)
        indx(i) = indx(j)
        indx(j)=itemp
        goto 3
5
        indx(l) = indx(j)
        indx(j) = indxt
        jstack=jstack+2
        if(jstack.gt.NSTACK)pause 'NSTACK too small in indexx'
        if(ir-i+1.ge.j-1)then
          istack(jstack)=ir
          istack(jstack-1)=i
          ir=j-1
        else
          istack(jstack)=j-1
          istack(jstack-1)=l
          l=i
        endif
      endif
      goto 1
      END
```

```
C
      Subroutine for initializing input parameters to CREME96 environment
C
C
      model.
C
      Modified 9/12/96: Energy range fixed at 0.1-1.0E+5 MeV/nuc;
C
                         However, the external flux routines return 0 for
C
                         E < 1.0 \text{ MeV/nuc}; the 0.1 threshold is put in here
С
                         for subsequent tracking through shielding.
C
C
C
      Modified 11/18/97: Allow input of trapped proton file.
С
C
C
       IMPLICIT NONE
       INTEGER*4 IMINTEMP, IMAXTEMP, IACCEPT, IFILETYPE
       INTEGER*4 IZMIN, IZMAX, IMODE, ITRANS, ITYPE, ITRP
       INTEGER*4 ISEPMODE
       REAL*4 EMIN, EMAX, YEAR, YEARDUM
       REAL*4 EMINTEMP, EMAXTEMP
       CHARACTER*80 GTRANSFILE, TRAPDFILE, FLXFILE
       CHARACTER*1 IBLANK
       DATA IBLANK/' '/
       INTEGER*4 IERR
       DATA IERR/0/
С
       WRITE(6,1000)
 1000 FORMAT(' CREME96 IONIZING RADIATION ENVIRONMENT MODEL',/,
             ' ----> FLUX DRIVER Module: External Environment')
       WRITE (6, 1001)
 1001 FORMAT (
     & 'This program will calculate the particle environment',

    ' outside of the spacecraft.',/,
        ' You must run additional programs after this to',
     & '(1) transport the particles'
     & /,' through shielding; and (2) calculate SEU rates.',
     &//,' BEFORE RUNNING THIS OR ANY OTHER CREME96 PROGRAM',
     & ' PLEASE DEFINE THREE LOGICALS:',/,
     & /,4x,' CREME96 as the directory where CREME96 source',
              ' & executables reside.',
      & /,4x,' CR96TABLES as the directory in which CREME96 data',
              ' tables reside.',
                             as the directory in which output files',
      & /,4x,' USER
              ' should be written.',
      & //,' Now begin specification of the environment parameters: ')
   101 CONTINUE
        CALL RETRY INPUT (IERR)
        WRITE(6,1002)
  1002 FORMAT(/,' Enter minimum & maximum atomic numbers: ',
        /,' Recommended for most applications: ',
           ' IZMIN = 1 (hydrogen) to IZMAX = 28 (nickel).',
          /,' [Enter 0 0 <CARRIAGE RETURN> for recommended values.] ',
                NOTE: For >95% of all SEU applications, Z > 28',
          ' elements, which are very',
         /,' rare, may be neglected. However, for SEU rates in',
      &
           ' devices with high',
```

```
thresholds (> 15 MeV-cm2/mg) these heavier elements',
     &
          ' MAY be important,',
     &
        /,' particularly for low-inclination low-Earth orbits',
     &
     &
         ' or for applications',
       /,' demanding very low SEU rates. Please note that',
        ' including Z > 28 elements in',
     &
        /,' you calculations can',
     &
          ' significantly slow down some parts of the CREME96 code.')
      READ (*, *, ERR=101, IOSTAT=IERR) IMINTEMP, IMAXTEMP
      IZMAX=MAX(IMINTEMP, IMAXTEMP)
       IZMIN=MIN(IMINTEMP, IMAXTEMP)
       IF (IZMIN.EQ.O .and. IZMAX.EQ.O) THEN
          TZMTN=1
          IZMAX=28
      ENDIF
      IF (IZMIN.LE.O .or. IZMAX.GT.92) THEN
         WRITE(6,9001) IZMIN, IZMAX
          FORMAT(1x,' Invalid atomic number(s): ',215,
 9001
              /,lx,' Please try again.')
          GOTO 101
      ENDIF
      WRITE(6,9002) IZMIN, IZMAX
 9002 FORMAT(1x,' Lowest atomic number = ', I5,
   & /,lx,' Highest atomic number = ',I5)
 103 CONTINUE
      EMIN=0.1
      EMAX=1.0E+5
C-----
C
     Following code for specifying energy interval obsolete 9/12/96:
C
      CALL RETRY INPUT (IERR)
С
      WRITE(6,1003)
C 1003 FORMAT(/,' Enter minimum & maximum energy (in MeV/nuc): ',
   & /,' Recommended for most SEE applications: ',
С
              ' EMIN = 10.0; EMAX = 1.0E+5')
C
С
      TYPE *,' [Enter 0 0 < CARRIAGE RETURN> for recommended values.]'
C
      READ(*,*,ERR=103,IOSTAT=IERR) EMINTEMP, EMAXTEMP
C
      EMIN=MIN (EMINTEMP, EMAXTEMP)
C
      EMAX=MAX (EMINTEMP, EMAXTEMP)
С
      IF (EMAX.LE.0.0 .and. EMIN.LE.0.0) THEN
C
       ENDIF
С
       IF (EMIN.LE.O .or. EMAX.LE.O.O .or. EMIN.EQ.EMAX) THEN
С
           TYPE *,' Invalid energy value(s): ',EMIN,EMAX
           TYPE *,' Please try again.'
C
C
           GOTO 103
C
       ENDIF
С
       TYPE *,' Minimum energy = ',EMIN,' MeV/nuc'
C
       TYPE *,' Maximum energy = ',EMAX,' MeV/nuc'
 104 CONTINUE
      CALL RETRY INPUT (IERR)
      WRITE (6, 1004)
 1004 FORMAT(/,' Specify type of environment model: ',
             ' Enter 0 or 1: ',
        /,' 0 = Solar-quiet (ie., no Solar Energetic Particles)',
    &
    & /,' 1 = Solar Energetic Particles ONLY')
```

```
WRITE (6, 1041)
                NOTE: Choosing 1 (Solar Energetic Particles ONLY)',
1041 FORMAT('
            ' does not include',
   &
          /,' Galactic cosmic rays, which may also contribute',
    &
           ' to the SEU rate behind',
          /,' thick shielding during a solar particle event.')
     READ (*, *, ERR=104, IOSTAT=IERR) ITYPE
     IF (ITYPE.NE.O .and. ITYPE.NE.1) THEN
         WRITE(6,9010) ITYPE
         FORMAT(1x,' Environment type ', I6,' unknown.',
9010
             /,1x,' Please try again.')
    æ
         GOTO 104
     ENDIF
     GTRANSFILE='
     TRAPDFILE ='
     CONTINUE
105
     CALL RETRY INPUT (IERR)
     IF (ITYPE.EQ.0) THEN
     IMODE=0
     WRITE (6, 1005)
1005 FORMAT(/,' Solar-quiet period. Enter decimal year',
   & '(eg. 1996.42) OR',
    & /,3x,'0 for Solar Minimum (Cosmic-Ray Maximum, YEAR =1977.0)',
    & /,3x,'1 For Solar Maximum (Cosmic-Ray Minimum, YEAR =1990.2)')
     READ(*,*,ERR=105,IOSTAT=IERR) YEARDUM
                         .LE. 0.0001) THEN
     IF (ABS (YEARDUM)
      YEAR=1977.0
      WRITE(6,9020) YEAR
9020 FORMAT(1x,' Solar Minimum (Cosmic-Ray Maximum) YEAR = ',F10.3)
     ELSEIF (ABS (YEARDUM-1.0) .LE. 0.0001) THEN
      YEAR=1990.2
      WRITE(6,9021) YEAR
9021
     FORMAT(1x,' Solar Maximum (Cosmic-Ray Minimum) YEAR = ',F10.3)
     ELSE
      YEAR=YEARDUM
      WRITE(6,9022) YEAR
     FORMAT(1x,' YEAR = ',F10.3)
9022
     ENDIF
     ELSEIF (ITYPE.EQ.1) THEN
     YEAR=0.0
 106 CONTINUE
     CALL RETRY INPUT (IERR)
     WRITE(6,1006)
1006 FORMAT(/,' CREME96 currently provides three Solar Energetic',
    &
            ' Particle Models: ',
    & /,3x,' Worst Week in 22 years: based on observed proton and',
   &
        ' heavy-ion fluences',
    & /,28x,' on 19-26 October 1989;',
    & /,3x,' Worst Day in 22 years: based on observed proton and',
            ' heavy-ion fluences',
       /,28x,' on 20 October 1989',
```

```
/,3x,' Peak Instantaneous Flux: based on peak 5-minute-'
     &
              'average fluxes observed',
     & /,28x,' during 19-26 October 1989.',
        /,3x,' Enter 1 for worst week; 2 for worst day;',
              ' 3 for peak flux: ')
       READ (*, *, ERR=106, IOSTAT=IERR) ISEPMODE
C
       Sloppy coding, introduced here on 9/14/96: ISEPMODE gives
С
       natural progression toward increasing severity, which is
C
C
       incompatible with original definitions of IMODE. Unfortunately,
       the IMODE values are deeply imbedded in the code and I have
C
С
       chosen not to change them at this time.
C
       IF (ISEPMODE.EQ.2) THEN
           IMODE=1
           WRITE(6,1007)
           FORMAT(' Worst Day Solar Energetic Particle Model chosen.')
 1007
       ELSEIF (ISEPMODE.EQ.1) THEN
           IMODE=2
           WRITE (6, 1008)
           FORMAT(' Worst Week Solar Energetic Particle Model chosen.')
 1008
       ELSEIF (ISEPMODE.EQ.3) THEN
           IMODE=3
           WRITE (6, 1081)
           FORMAT (' Peak Solar Energetic Particle Flux Model chosen.')
 1081
       ELSE
           WRITE(6,1009)
 1009
           FORMAT(' Requested SEP environment not defined.',
                  ' Please try again.')
           GOTO 106
       ENDIF
       ENDIF
 107 CONTINUE
       CALL RETRY INPUT (IERR)
       WRITE(6,1010)
 1010 FORMAT(/,' Specify Environment Location: ',
                ' Enter 0 or 1: ',
     ₽.
              /,' 0 = Interplanetary Space near Earth',
     &
                       (ie., outside of Earths magnetosphere)'
     &
              /,' 1 = Inside Earths magnetosphere',
                (You will need to supply a geomagnetic transmission',
     & /,'
               ' function file.',
     &
     & /,'
                 Run GTRANS DRIVER to make one.)')
       READ(*,*,ERR=107,IOSTAT=IERR) ITRANS
       IF (ITRANS.NE.O .and. ITRANS.NE.1) THEN
           WRITE(6,9030) ITRANS
 9030
           FORMAT(1x,' Environment location', I5,' unknown.',
     æ
                  ' Please try again.')
           GOTO 107
       ENDIF
       IF (ITRANS.EQ.0) THEN
           WRITE (6, 1011)
 1011
           FORMAT(' Geosynchronous Orbit or'
                  ' Near-Earth Interplanetary Space')
       ELSEIF (ITRANS.EQ.1) THEN
           CONTINUE
  112
```

```
CALL RETRY INPUT (IERR)
          WRITE (6, 1012)
          FORMAT(' Inside Earths Magnetosphere')
 1012
          WRITE (6, 1013)
          FORMAT(' Specify name of geomagnetic transmission file: ',
 1013
               /,' ie., something.GTF')
    &
          READ (*, 1014, ERR=112, IOSTAT=IERR) GTRANSFILE
          FORMAT (A80)
 1014
          IF (GTRANSFILE.EQ.IBLANK) THEN
          WRITE(6,1914)
           FORMAT(1x,' You must specify a geomagnetic transmission',
 1914
                     ' file (.GTF) for the calculation you',
    &
                /,1x,' have outlined. Please try again.',/)
    &
           GOTO 112
          ELSE
           IFILETYPE=2
           WRITE(6,1015) GTRANSFILE
           FORMAT(' Geomagnetic Transmission File =',/,1x,A80)
 1015
           CALL CHECK FILE (IFILETYPE, GTRANSFILE, IACCEPT)
           IF (IACCEPT.NE.0) GOTO 112
          ENDIF
          ITRP=0
          IF (IZMIN.EQ.1) THEN
          IF (IMODE.EQ.0) THEN
  116
          CONTINUE
          CALL RETRY_INPUT(IERR)
          WRITE(6,1016)
          FORMAT(' Include Trapped Protons? (0=no; 1=yes)')
 1016
           WRITE(6,9999)
C
           FORMAT(1x,' *** NOTE: This test version of CREME96',
C 9999
                    ' does NOT include trapped protons.',
C
                /,1x,' Please enter 0')
C
          READ(*,*,ERR=116,IOSTAT=IERR) ITRP
          ENDIF
          IF (ITRP.EQ.0) THEN
              WRITE(6,1027)
              FORMAT(1x,' No Trapped Protons Included.')
 1027
          ELSEIF (ITRP.EQ.1) THEN
              CONTINUE
  117
              CALL RETRY INPUT (IERR)
              WRITE(6,1017)
 1017
              FORMAT (' Trapped Protons included.',
                /,lx,' Enter name of',
    ۶
     &
              ' file containing orbit-averaged trapped proton flux:')
              READ(*,1014,ERR=117,IOSTAT=IERR) TRAPDFILE
              IF (TRAPDFILE.EQ.IBLANK) THEN
                 WRITE(6,1917)
 1917
                 FORMAT(1x,' You must specify a trapped proton',
                  ' file for the calculation you have outlined.',
    &
                  /,1x,' Please try again.',/)
     &
                 GOTO 117
              ELSE
```

```
IFILETYPE=1
                 WRITE(6,1018) TRAPDFILE
                 FORMAT(' Trapped Proton Flux File =',/,1x,A80)
1018
                 CALL CHECK_FILE(IFILETYPE, TRAPDFILE, IACCEPT)
                 IF (IACCEPT.NE.0) GOTO 117
                 ITRANS=2
              ENDIF
          ENDIF
          ENDIF
          ENDIF
          CONTINUE
 119
          CALL RETRY INPUT (IERR)
          WRITE(6,1019)
          FORMAT(/,' Particle environment specification now completed.',
1019
                 /,' Enter name of output file: '
    &
                 /,' Note: for standard CREME96 format, must be',
    &
                   ' something.FLX')
    &
          CONTINUE
120
          READ(*,1014,ERR=119,IOSTAT=IERR) FLXFILE
          WRITE(6,1020) FLXFILE
              FORMAT(' Output FLux File =',/,1x,A80)
1020
          CALL CHECK OUTPUT FILE (FLXFILE, IACCEPT)
          IF (IACCEPT.NE.0) GOTO 120
          RETURN
          END
```

```
SUBROUTINE INIDOSE (INFILE, LETMINMG, LETMAXMG,
                         IZMIN, IZMAX, EMINCUT, EMAXCUT, MATERIAL, OUTFILE)
     &
C
       Subroutine for initializing input parameters to the DOSE program
C
       in CREME96. This version only allows SILICON devices.
C
C
       IMPLICIT NONE
       INTEGER*4 IZMIN, IZMAX, IMINTEMP, IMAXTEMP, IFILETYPE, IACCEPT
       REAL*4 LETMINMG, LETMAXMG, LETMINTEMP, LETMAXTEMP
       REAL*4 EMINCUT, EMAXCUT
       CHARACTER*80 INFILE, OUTFILE, DEFAULT NAME
       CHARACTER*12 MATERIAL
       CHARACTER*1 IBLANK
       DATA IBLANK/' '/
       INTEGER*4 IERR, IDIFSPEC, ILONG
       DATA IERR/0/
C
       WRITE(6,1000)
 1000 FORMAT (' CREME96 IONIZING RADIATION ENVIRONMENT MODEL',
     & /,' --> Ionizing Dose Calculation')
       WRITE(6,1001)
 1001 FORMAT(' This program will calculate the dose resulting from',
     & 'CREME96 differential',
     & /,' particle fluxes. This program is intended primarily for',
     & ' calculating dose due to',
     & /,' NON-TRAPPED components of the radiation environment',
     & ' [cosmic rays and solar',/,' energetic',
        ' (flare) particles].',
     &
       ' This program is NOT recommended for calculating',
     & /,' dose due to TRAPPED particles,',
     & ' which generally dominate the dose inside',
     & /,' Earths magnetosphere.',
     & 'CREME96 does NOT included trapped electrons, and',
     & /,' trapped-proton dose is more accurately described',
     & ' by other programs, especially',
     & /,' for lightly-shielded systems.',
     &//,' Before running this program, you must do:',
     &//,' FLUX',
         ' to generate the particle environment outside',
     & ' the spacecraft; and',
     & /,' TRANS',
     & ' to transport fluxes through the spacecraft shielding.',
     &//,' NOTE: Before running this or any other CREME96 program',
     & ' please define 3 logicals:',/,
                            as the directory where CREME96 source',
     & /,4x,' CREME96
              ' & executables reside.',
     & /,4x,' CR96TABLES as the directory in which CREME96 data',
              ' tables reside.',
                            as the directory in which output files',
     & /,4x,' USER
              ' should be written.',
     & //,' Now begin specification of the DOSE_DRIVER inputs: ')
        INFILE='
  112 CONTINUE
       CALL RETRY INPUT (IERR)
       WRITE(6,1002)
  1002 FORMAT(' Enter name of file containing',
                ' CREME96 particle fluxes: '
      æ
               /,' ie. something.TFX from TRANS or',
```

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```
(for zero shielding) something.FLX from FLUX'
               ,' or something.TR* from TRP:')
       READ(*,1014,ERR=112,IOSTAT=IERR) INFILE
 1014 FORMAT (A80)
       IF (INFILE.EQ.IBLANK) THEN
           WRITE(6,1914)
          FORMAT(1x,' You must specify an input file here,',
 1914
                     ' either from TRANS or',
     æ
           /,lx,' (in the case of zero shielding) from',
                ' FLUX or TRP. ',
     ۶
                     ' Please try again.',/)
            GOTO 112
       ELSE
           IFILETYPE=4
          WRITE(6,1020) INFILE
          FORMAT(' Input Flux File =',/,1x,A80)
 1020
           CALL CHECK FILE (IFILETYPE, INFILE, IACCEPT)
           IF (IACCEPT.NE.0) GOTO 112
       ENDIF
  103 CONTINUE
C
C
       Modification 9/12/96: LET range hardwired:
       LETMINMG=1.0E-3
       LETMAXMG=1.1E+2
1032 CONTINUE
       CALL RETRY INPUT (IERR)
       WRITE(6,1004)
 1004 FORMAT(' Enter minimum & maximum atomic numbers to be',
                ' included in dose calculation:',
     &
                   [Enter 0 0 < CARRIAGE RETURN> for full range',
     &
                ' included in the input flux file.]')
     &
      CONTINUE
       READ(*,*,ERR=1032,IOSTAT=IERR) IMINTEMP,IMAXTEMP
       IF (IMAXTEMP.NE.O .and. IMAXTEMP.LT.IMINTEMP) THEN
           IZMIN=MIN(IMINTEMP, IMAXTEMP)
           IZMAX=MAX(IMINTEMP,IMAXTEMP)
       ELSE
           IZMIN=IMINTEMP
           IZMAX=IMAXTEMP
       ENDIF
       IF (IZMIN.LT.0 .or. IZMIN.GT.92
           .or. IZMAX.LT.0 .or. IZMAX.GT.92) THEN
               WRITE(6,9002) IZMIN, IZMAX
 9002
              FORMAT(1x,' Invalid atomic number(s): ',215,
                    /,1x,' Please try again.')
     &
               GOTO 101
       ENDIF
       IF (IZMIN.EQ.O .and. IZMAX.EQ.O) THEN
           WRITE(6,1039)
 1039 FORMAT(' Nominal Z range from input flux file used.')
       ELSEIF (IZMIN.EQ.O .and. IZMAX.NE.O) THEN
           WRITE(6,1038) IZMAX
 1038 FORMAT(' Minimum Z value from input flux file; Maximum Z =',I3)
```

```
ELSEIF (IZMIN.NE.O .and. IZMAX.EQ.O) THEN
             WRITE(6,1037) IZMIN
    1037 FORMAT(' Minimum Z =', I3,'; Maximum Z value from',
       & 'input flux file.')
         ELSE
             WRITE(6,1040) IZMIN, IZMAX
    1040 FORMAT(1x,' Dose accumulated for elements',
       \& /,1x,13,' </= Z </= ',13)
         ENDIF
         12/1/97: EMIN, EMAX hardwired. Keep source code here in case requested
   С
        by beta-testers.
   C
         EMINCUT=0.1
         EMAXCUT=1.0E+5
   C-----
   C 105 CONTINUE
          CALL RETRY INPUT (IERR)
   C
          EMAXCUT=1.0E+24
   C
         WRITE(6,1005)
   C 1005 FORMAT(' Enter minimum particle energy (in MeV/nuc)',
   C & 'to be included in accumulating the'
              /,' dose calculation:')
   C
   C
          READ(*,*,ERR=105,IOSTAT=IERR) EMINCUT
   C
          IF (EMINCUT.LT.O.) THEN
  C
  C
             WRITE(6,9005) EMINCUT
              FORMAT(1x,' Invalid minimum energy value: ',E13.6,
  C 9005
             /,lx,' Please try again.')
   C
              GOTO 105
   С
   C
          ENDIF
         WRITE(6,1051) EMINCUT
   C
   C 1051 FORMAT(/,' Dose accumulated for',
        % ' nuclei with energy > ',F8.3,' MeV/nuc.')
Brid.
          MATERIAL='SILICON'
          WRITE(6,1007) MATERIAL
    1007 FORMAT(/,' Dose calculated in ',A12)
    1017 CONTINUE
          CALL RETRY INPUT (IERR)
          WRITE(6,1008)
    1008 FORMAT(/,' Enter name of output file: ',
        & /,' Note: According to CREME96 naming conventions,',
                ' should be something.dse')
        &
          ILONG=INDEX (INFILE, '.')
          IF (ILONG.NE.0) THEN
             DEFAULT_NAME=INFILE(1:ILONG)//'DSE'
          ELSE
             DEFAULT_NAME=INFILE//'.DSE'
          ENDIF
          WRITE(6,1028) DEFAULT_NAME(1:79)
     1028 FORMAT (' Suggested name: ', /, 1x, A79,
          /,' Hit RETURN if this is acceptable.')
```

1018 CONTINUE

```
READ(*,1014,ERR=1017,IOSTAT=IERR) OUTFILE

IF (OUTFILE.EQ.IBLANK) OUTFILE=DEFAULT_NAME

WRITE(6,1009) OUTFILE

1009 FORMAT(' Output Flux File =',/,lx,A80)

CALL CHECK_NAME_CONFLICT(INFILE,OUTFILE,IACCEPT)

IF (IACCEPT.NE.0) GOTO 1017

CALL CHECK_OUTPUT_FILE(OUTFILE,IACCEPT)

IF (IACCEPT.NE.0) THEN

WRITE(6,1010) INFILE(1:75)

WRITE(6,1011) OUTFILE(1:75)

WRITE(6,1012)

1010 FORMAT(1x,' INPUT file = ',/,5x,A75)

1011 FORMAT(1x,' Previous try at OUTPUT name = ',/,5x,A75)

1012 FORMAT(1x,' Try again, ie., newname:DSE')

GOTO 1018

ENDIF
```

RETURN END

```
SUBROUTINE INILET (INFILE, LETMINMG, LETMAXMG,
     ۶
                          IZMIN, IZMAX, EMINCUT, EMAXCUT, MATERIAL, OUTFILE,
     æ
                          IDIFSPEC)
C
C
       Subroutine for initializing input parameters to LETSPEC program
C
       in CREME96. This version only allows SILICON devices.
C
C
       Modifications 11/8/96: Allow 0 for input IZ values to select
C
                               either lowest or highest from input flux
C
                               file.
C
C
       Modification 10/31/96: option of differential LET spectrum added.
C
C
       Modifications 9/12/96: LETMINMG, LETMAXMG hardwired;
C
                               allow user to specify minimum energy in
C
                              flux accumulation.
C
       IMPLICIT NONE
       INTEGER*4 IZMIN,IZMAX,IMINTEMP,IMAXTEMP,IFILETYPE,IACCEPT
       REAL*4 LETMINMG, LETMAXMG, LETMINTEMP, LETMAXTEMP
       REAL*4 EMINCUT, EMAXCUT
       CHARACTER*80 INFILE, OUTFILE, DEFAULT NAME
       CHARACTER*12 MATERIAL
       CHARACTER*1 IBLANK
       DATA IBLANK/' '/
       INTEGER*4 IERR, IDIFSPEC, ILONG
       DATA IERR/0/
       WRITE(6,1000)
 1000 FORMAT (' CREME96 IONIZING RADIATION ENVIRONMENT MODEL',
     & /,' --> INTEGRAL Linear Energy Transfer (LET)',
     & ' Spectrum Calculation')
       WRITE(6,1001)
 1001 FORMAT(' This program will transform the input differential',
     & ' particle energy spectra from',
     & /,' CREME96 into an LET spectrum, ie.,',
     & ' particle flux vs. LET [in MeV-cm2/g]',
     & /,' as appropriate for SEU calculations with CREME96. Before',
     & 'running this',
     & /,' program, you must do:',
            FLUX ',
     &//,'
         ' to generate the particle'
       ' environment outside the spacecraft; and'
             TRANS',
         ' to transport fluxes through the spacecraft shielding.',
     &//,' NOTE: Before running this or any other CREME96 programs',
     & ' please define three ',
     & /,' logicals: ',/,
     & /,4x,' CREME96
                            as the directory where CREME96 source',
              ' & executables reside.',
     & /,4x,' CR96TABLES
                           as the directory in which CREME96 data',
              ' tables reside.',
     & /,4x,' USER
                            as the directory in which output files',
              ' should be written.',
     & //,' Now begin specification of the LETSPEC DRIVER inputs: ')
       INFILE='
```

```
CALL RETRY INPUT (IERR)
        WRITE(6,1002)
 1002 FORMAT(' Enter name of file containing',
                ' CREME96 particle fluxes, ie. something.TFX',
               /,' from TRANS or',
     &
                ' (for zero shielding) something.FLX from FLUX'
      3
               /,' or something.tr* from TRP:')
       READ (*, 1014, ERR=112, IOSTAT=IERR) INFILE
 1014 FORMAT (A80)
       IF (INFILE.EQ.IBLANK) THEN
            WRITE(6,1914)
            FORMAT(1x,' You must specify an input .FLX file here,',
 1914
                     ' either from TRANS or',
           /,lx,' (in the case of zero shielding) from',
                 ' FLUX. ',
                     ' Please try again.',/)
       ELSE
            IFILETYPE=4
            WRITE(6,1020) INFILE
            FORMAT(' Input Flux File =',/,1x,A80)
 1020
            CALL CHECK FILE (IFILETYPE, INFILE, IACCEPT)
            IF (IACCEPT.NE.0) GOTO 112
       ENDIF
  103 CONTINUE
С
       Modification 9/12/96: LET range hardwired:
           LETMINMG=1.0E-3
           LETMAXMG=1.1E+2
C
     Following code obsolete 9/12/96:
C
C
    WRITE(6,1003)
C 1003 FORMAT(/,' Enter minimum & maximum LET values (in MeV-cm2/mg)',
              /,' [Recommended for most SEE applications:'
С
      &
               /,' minimum LET = 1.0E-3 MeV-cm2/mg ',
/,' maximum LET = 1.1E+2 MeV-cm2/mg] ',
C
               /,'
C
               /,' NOTE THE UNITS USED HERE -- per milligram!',
С
      & /,' Enter 0 0 <CARRIAGE RETURN> for recommended defaults.)')
C
C
       ACCEPT *, LETMINTEMP, LETMAXTEMP
C .
       LETMINMG=MIN(LETMINTEMP, LETMAXTEMP)
C
        LETMAXMG=MAX (LETMINTEMP, LETMAXTEMP)
C
       IF ((LETMINMG.EQ.O. .and. LETMAXMG.EQ.O.)
C
            .or. (LETMAXMG.LE.LETMINMG) ) THEN
C
            LETMINMG=1.0E-3
C
            LETMAXMG=1.1E+2
C
       ENDIF
C
       IF (LETMINMG.LT.O. .or. LETMAXMG.LT.O.
С
            .or.LETMINMG.EQ.LETMAXMG) THEN
С
           WRITE (6,9001) LETMINMG, LETMAXMG
C 9001
          FORMAT(1x,' Invalid LET value(s): ',E13.6,2x,E13.6,
C
                /,1x,' Please try again.')
C
           GOTO 103
C ·
       ENDIF
        WRITE(6,1031) LETMINMG, LETMAXMG
 1031 FORMAT(/,' Integral LET spectrum accumulated for ',
               /,1x,E12.5,' </= LET </= ',E12.5,' MeV-cm2/mg',/)
```

```
1032 CONTINUE
      CALL RETRY INPUT (IERR)
      WRITE (6, 1004)
1004 FORMAT(' Enter minimum & maximum atomic numbers to be',
               ' included in integral LET spectrum:',
                  [Enter 0 0 <CARRIAGE RETURN> for full range',
    &
               ' included in the input flux file.',
    æ
              /,' NOTE: in general',
                ' protons (Z=1) should NOT be included in the',
                ' LET spectrum',/,' for most SEU calculations.]')
    CONTINUE
101
      READ(*,*,ERR=1032,IOSTAT=IERR) IMINTEMP,IMAXTEMP
      IF (IMAXTEMP.NE.O .and. IMAXTEMP.LT.IMINTEMP) THEN
          IZMIN=MIN(IMINTEMP, IMAXTEMP)
          IZMAX=MAX(IMINTEMP,IMAXTEMP)
      FLSE
          IZMIN=IMINTEMP
          IZMAX=IMAXTEMP
      ENDIF
      IF (IZMIN.LT.0 .or. IZMIN.GT.92
         .or. IZMAX.LT.0 .or. IZMAX.GT.92) THEN
              WRITE(6,9002) IZMIN, IZMAX
             FORMAT(1x,' Invalid atomic number(s): ',215,
9002
                   /,1x,' Please try again.')
              GOTO 101
      ENDIF
      IF (IZMIN.EQ.O .and. IZMAX.EQ.O) THEN
          WRITE(6,1039)
1039 FORMAT(' Nominal Z range from input flux file used.',/)
      ELSEIF (IZMIN.EQ.0 .and. IZMAX.NE.0) THEN
          WRITE(6,1038) IZMAX
1038 FORMAT(' Minimum Z value from input flux file; Maximum Z =',I3)
      ELSEIF (IZMIN.NE.0 .and. IZMAX.EQ.0) THEN
          WRITE(6,1037) IZMIN
1037 FORMAT(' Minimum Z =', I3,'; Maximum Z value from',
            ' input flux file.')
    ٠. ي
      ELSE
          WRITE(6,1040) IZMIN, IZMAX
1040 FORMAT(1x,' Integral LET spectrum accumulated for elements',
    \& /,1x,13,' </= Z </= ',13,/)
      ENDIF
 105 CONTINUE
      CALL RETRY INPUT (IERR)
      EMAXCUT=1.0E+24
      WRITE(6,1005)
1005 FORMAT(' Enter minimum particle energy (in MeV/nuc)',
             ' to be included in accumulating the'
    &
           /,' integral LET spectrum:',
              [NOTE: for most SEU applications,',
    æ
          ' the recommended value = 0.1 MeV/nuc.',
          /,' However, in some devices, ranging out of low-energy',
    &
          ' particles along very',
    &
                long RPP chords can lead to gross overestimates, ',
    &
           ' particularly for low-',
```

```
/,' threshold devices in solar particle events. In',
          ' these cases, larger minimum',
               energy values (1-10 MeV/nuc) should be explored.]')
       READ(*,*,ERR=105,IOSTAT=IERR) EMINCUT
       IF (EMINCUT.LT.O.) THEN
          WRITE(6,9005) EMINCUT
          FORMAT(1x,' Invalid minimum energy value: ',E13.6,
9005
             /,1x,' Please try again.')
          GOTO 105
       ENDIF
     WRITE (6, 1051) EMINCUT
1051 FORMAT(/,' Integral LET spectrum accumulated for',
              ' nuclei with energy > ',F8.3,' MeV/nuc.')
     MATERIAL='SILICON'
     WRITE (6, 1007) MATERIAL
1007 FORMAT(/,' LET spectrum calculated in ',A12)
1017 CONTINUE
     CALL RETRY INPUT (IERR)
     WRITE (6, 1008)
1008 FORMAT(/,' Enter name of output file: ',
            /,' Note: for standard input to CREME96 SEU routines',
    &
             ' must be something.LET')
     ILONG=INDEX(INFILE, '.')
     IF (ILONG.NE.O) THEN
         DEFAULT NAME=INFILE(1:ILONG)//'LET'
          DEFAULT NAME=INFILE//'.LET'
      ENDIF
     WRITE(6,1028) DEFAULT_NAME(1:79)
1028 FORMAT(' Suggested name:',/,1x,A79,
           /,' Hit RETURN if this is acceptable.')
1018 CONTINUE
     READ (*, 1014, ERR=1017, IOSTAT=IERR) OUTFILE
      IF (OUTFILE.EQ.IBLANK) OUTFILE=DEFAULT NAME
     WRITE(6,1009) OUTFILE
1009 FORMAT(' Output Flux File =',/,1x,A80)
      CALL CHECK NAME CONFLICT (INFILE, OUTFILE, IACCEPT)
      IF (IACCEPT.NE.0) GOTO 1017
     CALL CHECK OUTPUT FILE (OUTFILE, IACCEPT)
     IF (IACCEPT.NE.0) THEN
     WRITE(6,1010) INFILE(1:75)
     WRITE (6, 1011) OUTFILE (1:75)
     WRITE(6,1012)
1010 FORMAT(1x,' INPUT file = ',/,5x,A75)
1011 FORMAT(1x,' Previous try at OUTPUT name = ',/,5x,A75)
1012 FORMAT(1x,' Try again, ie., newname.LET')
     GOTO 1018
      ENDIF
```

```
1060 CONTINUE
     CALL RETRY INPUT (IERR)
     WRITE(6,1061)
1061 FORMAT(/,' Do you want a DIFFERENTIAL LET spectrum also?: ',
              ' (0=no; 1=yes) ',
           /,' NOTE: A differential LET spectrum is NOT necessary',
    &
           ' for SEU calculations.')
     READ(*, *, ERR=1060, IOSTAT=IERR) IDIFSPEC
     IF (IDIFSPEC.NE.1) IDIFSPEC=0
     IF (IDIFSPEC.EQ.0) WRITE(6,1062)
     IF (IDIFSPEC.EQ.1) WRITE(6,1063)
1062 FORMAT(' No differential LET spectrum will be created.',/)
1063 FORMAT(' Differential LET spectrum also created. The file name',
          ' will be the same as',
          /,' that of the integral LET spectrum,'
    &
           ' but with extension .DLT',/)
```

RETURN END

WRITE(6,1002)

æ

```
C
       Subroutine for initializing input parameters to transport routine
C
       in CREME96. This version only allows ALUMINUM shielding.
С
С
      Modified 06-13-96: to include shielding distribution
C
      Modified 11-13-96: gets shielding distribution from standard *.SHD
C
                          file, as created with the SHIELDFILE_DRIVER program.
C
      Modified 11-17-97: to include .trp inputs
C
C
       IMPLICIT NONE
       INTEGER*4 IPATH, IULABEL, IFILETYPE, IACCEPT
       INTEGER*4 KFILETYPE, KACCEPT
       REAL*4 UPATH
       CHARACTER*1 IBLANK
       DATA IBLANK/' '/
       CHARACTER*80 INFILE, SHIELDFILE, OUTFILE
       CHARACTER*12 TARGET
       CHARACTER*5 UNITS LABEL
       DIMENSION UNITS LABEL(4)
       DATA UNITS_LABEL/'g/cm2','mils ','cm ','!!!!!'/
       INTEGER*4 IERR
       DATA IERR/0/
С
      WRITE(6,1000)
 1000 FORMAT (' CREME96 IONIZING RADIATION ENVIRONMENT MODEL',
            /,' --> NUCLEAR TRANSPORT PROGRAM')
      WRITE(6,1001)
 1001 FORMAT(' This program will transport the ionizing',
             '-radiation particle fluxes generated by',
            /,' the CREME96 code through aluminum shielding',
     æ
            ' of specified thickness. Before',
           /,' running this program, you must do FLUX',
     &
             ' (ie, run CREME96:FLUX_DRIVER)',
     ۶
          /,' or TRP',
     &
              ' (ie, run CREME96:TRAPPED PROTON_DRIVER)',
             ' to generate', /, ' the particle environment',
             ' outside of the spacecraft. After running',
     &
              ' this program', /, ' you will run other routines to',
              ' calculate SEU rates. ',
     \epsilon //,' NOTE: Before running this or any other CREME96 programs',
     & ' please define three ',
     & /,' logicals: ',/,
                            as the directory where CREME96 source',
     & /,4x,' CREME96
              ' & executables reside.',
     & /,4x,' CR96TABLES as the directory in which CREME96 data',
              ' tables reside.',
                            as the directory in which output files',
     & /,4x,' USER
              ' should be written.',
     & //,' Now begin specification of the transport parameters: ')
       INFILE='
  112 CONTINUE
       CALL RETRY INPUT (IERR)
```

SUBROUTINE INIPROP(INFILE, IPATH, UPATH, TARGET,

SHIELDFILE, OUTFILE)

```
1002 FORMAT(/,' Enter name of file containing',
              ' CREME96 particle fluxes:',
    &
             /,' ie., something.FLX, something.TRP, or something.TFX')
     READ (*, 1014, ERR=112, IOSTAT=IERR) INFILE
1014 FORMAT (A80)
      IF (INFILE.EQ.IBLANK) THEN
          WRITE(6,1914)
          FORMAT(1x,' You must specify here EITHER a .FLX file',
1914
                    ' from FLUX (FLUX DRIVER)',
               /,1x,' OR a .TR* file from a ',
    &
                    ' TRP (TRAPPED PROTON DRIVER). ',
    &
               /,1x,' OR a .TFX file from a previous run of',
    æ
                    ' TRANS (TRANSPORT DRIVER). ',
    &
               /,1x,' Please try again.',/)
    &
           GOTO 112
      ELSE
          IFILETYPE=3
          WRITE(6,1020) INFILE
          FORMAT(' Input Flux File =',/,1x,A80)
1020
          CALL CHECK FILE (IFILETYPE, INFILE, IACCEPT)
          IF (IACCEPT.NE.0) GOTO 112
      ENDIF
      TARGET='ALUMINUM'
1021 CONTINUE
      CALL RETRY INPUT (IERR)
      WRITE(6,1003) TARGET(1:8)
1003 FORMAT(/,' In what units will the ',A8,
             ' shielding thickness be given? ',
   &
             /,' Enter 0, 1, or 2: ',
                                     0 = g/cm**2',
    &
             /,'
                                     1 = mils ',
            1,'
    &
                                     2 = cm
            /,'
    &
            //, (Note: 100 mils = 0.254 cm = 0.6858 g/cm**2 Al.)')
    &
      READ(*,*,ERR=1021,IOSTAT=IERR) IPATH
      IF (IPATH.LT.0 .or. IPATH.GT.2) THEN
          WRITE(6,9000)
          FORMAT(1x,' Illegal units specification. Please try again.')
9000
          GOTO 1021
      ENDIF
      IULABEL=IPATH+1
      IF (IULABEL.GT.4) IULABEL=4
      WRITE(6,9001) UNITS LABEL(IULABEL), TARGET
9001 FORMAT(1x,' Shielding thicknesses: in ',A5,1x,A12)
      SHIELDFILE='
      WRITE(6,1035)
1035 FORMAT(/,1x,' COMMENT ON SHIELDING VALUES: It is common',
                 ' practice for researchers dealing',
    €
             /,1x,' with total dose and dose-rate effects to',
    &
                 ' determine part response with zero',
    &
             /,1x,' shielding. For single event effects, on the '
    æ
                 ' other hand, it is important to',
    &
             /,1x,' shield out low-energy particles',
    &
                 ' which would never be encountered in a',
    &
             /,1x,' realistic situation. A nominal shielding',
    &
                   ' thickness of 100 mils is therefore',
```

```
/,1x,' recommended for general comparison purposes.',
     &
                   ' However, a realistic',
     æ
                   ' shielding',
     &
              /,1x,' distribution is essential for accurate SEU',
                   ' calculations in solar energetic',
              /,lx,' particle ("flare") environments.')
 1036 CONTINUE
       CALL RETRY INPUT (IERR)
       WRITE(6,1004)
 1004 FORMAT(/,1x,' Enter shielding thickness: ',
              /,1x' [Enter 0 if you wish to specify a file'
                  ' containing a shielding distribution.]')
     &
       READ (*, *, ERR=1036, IOSTAT=IERR) UPATH
       IF (UPATH.GT.0.0) THEN
           WRITE (6, 1005) UPATH, UNITS LABEL (IULABEL), TARGET
           FORMAT(' Shielding thickness = ',F10.5,1x,A5,5x,A12)
 1005
       ELSE
          CONTINUE
 1039
          CALL RETRY INPUT (IERR)
          WRITE (6, 1041)
 1041 FORMAT(/,lx,' Enter name of file containing shielding',
                ' distribution. This file, which should',
     &
           /,lx,' be called something.SHD, has a special',
     &
                ' format. This file must have been',
     &
            /,lx,' created before running TRANS with the',
     &
                 ' CREME96 command SHIELDFILE.')
     &
          READ (*, 1014, ERR=1039, IOSTAT=IERR) SHIELDFILE
          WRITE (6, 1042) SHIELDFILE
          FORMAT(' Shielding File =',/,1x,A80)
 1042
          KACCEPT=0
C
          KFILETYPE=0
          KFILETYPE=7
          CALL CHECK FILE (KFILETYPE, SHIELDFILE, KACCEPT)
          IF (KACCEPT.NE.0) GOTO 1039
       ENDIF
       WRITE(6,1008)
 1008 FORMAT(/,' Enter name of output file, ie., newname.TFX:')
 1043 CONTINUE
       CALL RETRY INPUT (IERR)
       READ(*,1014,ERR=1043,IOSTAT=IERR) OUTFILE
       WRITE(6,1009) OUTFILE
 1009 FORMAT(' Output Flux File =',/,1x,A80)
       CALL CHECK OUTPUT FILE (OUTFILE, IACCEPT)
       IF (IACCEPT.NE.O) THEN
       WRITE(6,1010) INFILE(1:75)
       WRITE(6,1011) OUTFILE(1:75)
       WRITE(6,1012)
 1010 FORMAT(1x,' INPUT file = ',/,5x,A75)
 1011 FORMAT(1x,' Previous try at OUTPUT name = ',/,5x,A75)
 1012 FORMAT(1x,' Try again, ie., newname.TFX')
       GOTO 1043
       ENDIF
```

The state of the s

```
IUNITS, MATERIAL, NBINS, XTHICK, XPROB,
         &
                                SHIELDFILE)
         &
   C
          Subroutine for initializing inputs to CREME96 program which
   C
           creates a shielding distribution file in standard format.
   C
          This version only allows ALUMINUM shielding.
   C
   С
           IMPLICIT NONE
           INTEGER*4 IUNITS, NBINS, MAXSHIELD, IULABEL
           CHARACTER*12 MATERIAL
           CHARACTER* 1 IBLANK
          DATA IBLANK/' '/
          REAL*4 XTHICK(1), XPROB(1)
          REAL*4 XTEMP, PTEMP
           CHARACTER*80 COMMENT, SHIELDFILE
          CHARACTER*5 UNITS LABEL
          DIMENSION UNITS LABEL (4)
          DATA UNITS LABEL/'g/cm2', 'mils ', 'cm ', '!!!!!'/
   C
          INTEGER*4 IERR, IACCEPT
          DATA IERR/0/
   C
          WRITE (6, 1000)
1000 FORMAT (' CREME96 IONIZING RADIATION ENVIRONMENT MODEL',
               /,' --> Shielding distribution program')
WRITE (6, 1001)
    1001 FORMAT(' This auxilliary program will use',
' specified inputs to create a shielding '
1
        &
                /,' distribution file with the format',
                 ' and header information expected by CREME96.',
Ŀ
         &//,' NOTE: Before running this program please define',
#
                 ' three logicals:',/,
Li
                          as the directory where CREME96 source',
         & /,4x,' CREME96
                  ' & executables reside.',
        &
         & /,4x,' CR96TABLES as the directory in which CREME96 data',
Ē.ā.
                 ' tables reside.',
111
                                as the directory in which output files',
        & /,4x,' USER
                 ' should be written.',
         & //,' Now begin specification of the shieldfile inputs: ')
     105
            CONTINUE
            CALL RETRY INPUT (IERR)
            WRITE(6,1215)
            FORMAT(/,1x,' Enter comment (80 characters max)',
    1215
                        ' for record-keeping in output file:')
            READ(*,1218,ERR=105,IOSTAT=IERR) COMMENT
           FORMAT (A80)
    1218
    1021 CONTINUE
           CALL RETRY INPUT (IERR)
           WRITE(6,1003)
    1003 FORMAT(/,' In what units will the shielding thickness be given?',
                   ' Enter 0, 1, or 2: ',
         &
                  1.
                          0 = g/cm**2',
                  1,'
                           1 = mils
                   /,'
                            2 = cm
```

/,' (Note: 100 mils = 0.254 cm = 0.6858 g/cm**2 Al.)')

SUBROUTINE INISHIELD (MAXSHIELD, COMMENT,

```
READ(*, *, ERR=1021, IOSTAT=IERR) IUNITS
      IF (IUNITS.LT.O .or. IUNITS.GT.2) THEN
          WRITE(6,9003)
          FORMAT(1x,' Illegal units specification. Please try again.')
9003
          GOTO 1021
      ENDIF
      IULABEL=IUNITS+1
      IF (IULABEL.GT.4) IULABEL=4
      MATERIAL='ALUMINUM'
      WRITE(6,9000) UNITS LABEL(IULABEL), MATERIAL
9000 FORMAT(1x,' Shielding thickness in', A5, 2x, A12)
      NBINS=0
      WRITE(6,1041) MAXSHIELD
1041 FORMAT(/,1x,' Now begin entry of shielding distribution: ',
    & /,1x,' Enter thickness (in the units specified above) and',
           ' coverage factor (a number',
    & /,1x,' between 0 and 1) then <CARRIAGE RETURN>',
    & ' for each bin of the distribution.',
    & /,1x,' Terminate your input list with 0 0 <CARRIAGE RETURN>.',
    & /,1x,' The maximum number of bins allowed in the',
          ' distribution is', I4,'.')
1036 CONTINUE
     WRITE(6,1042)
1042 FORMAT(/,1x,' Enter shielding thickness and coverage factor: ')
      CALL RETRY INPUT (IERR)
      READ(*, *, ERR=1036, IOSTAT=IERR) XTEMP, PTEMP
      IF (XTEMP.GT.0.0 .and. PTEMP.GT.0.0) THEN
           NBINS=NBINS+1
           IF (NBINS.LE.MAXSHIELD) THEN
               XTHICK (NBINS) = XTEMP
               XPROB (NBINS) = PTEMP
               WRITE(6,999) NBINS, XTHICK(NBINS), UNITS_LABEL(IULABEL),
                            XPROB (NBINS)
999
               FORMAT(1x,' SHIELDING BIN', 14,' THICKNESS = ',
                           F10.4,1x,A5,'
                                          FRACTION = ', F8.4
           ELSE
               WRITE(6,1043) NBINS, MAXSHIELD
1043
               FORMAT(1x,' Input terminated: No. input bins = ', 15,
                    /,1x,'
                                             Maximum allowed = ', I5)
               GOTO 1050
           ENDIF
      ELSE
           WRITE (6, 1044)
1044
           FORMAT(1x,' Shielding distribution input completed.')
           GOTO 1050
      ENDIF
      GOTO 1036
1050 CONTINUE
     SHIELDFILE='
 112 CONTINUE
      CALL RETRY INPUT (IERR)
```

```
WRITE(6,1002)
1002 FORMAT(/,' Enter name of output shielding file:',
              ' ie., something.SHD (Your file must have',
    &
             /,' this extension in order',
    &
               ' to be accessbile by CREME96',
    &
               ' directory routines.)')
      READ(*,1014,ERR=112,IOSTAT=IERR) SHIELDFILE
1014 FORMAT (A80)
      IF (SHIELDFILE.EQ.IBLANK) THEN
          WRITE(6,1914)
1914
          FORMAT(1x,' You must specify a filename here:',
               /,1x,' Please try again.',/)
    &
           GOTO 112
      ELSE
          WRITE(6,1020) ShieldFILE
1020
          FORMAT(' Output Shielding Distribution File =',/,lx,A80)
          CALL CHECK_OUTPUT_FILE(SHIELDFILE, IACCEPT)
          IF (IACCEPT.NE.0) GOTO 112
      ENDIF
      RETURN
     END
```

```
SUBROUTINE INITIALIZE HI UPSETS (LET FILE, XM, YM, ZM, FUNNEL, NBITS,
                                        IPARAM, PARAMS, XSECT FILE, IENTER,
     &
                                        DEVICE LABEL, REPORT FILE)
     &
C
C
C
        Generates interactive dialogue to get necessary input parameters
        for heavy-ion upsets:
С
C
        Written by:
                       Allan J. Tylka
C
                       Code 7654
С
                       Naval Research Laboratory
C
                       Washington, DC 20375-5352
C
                       tylka@crs2.nrl.navy.mil
С
C
С
C-
C
        IMPLICIT NONE
        CHARACTER*80 LET FILE, XSECT_FILE, REPORT_FILE
        CHARACTER*40 DEVICE LABEL
        REAL*4 XM, YM, ZM, FUNNEL, PARAMS, NBITS
        INTEGER*4 IPARAM, IACCEPT, IFILETYPE, IENTER
        CHARACTER*1 IBLANK
        DATA IBLANK/' '/
        DIMENSION PARAMS (4)
        INTEGER*4 IERR
       DATA IERR/0/
       IF (IENTER.EQ.1) THEN
       WRITE(6,1000)
 1000 FORMAT(' CREME96 IONIZING RADIATION ENVIRONMENT MODEL',
     & /,' --> Heavy-Ion-Induced Single Event Upset',
     & ' (SEU) Rate Calculation')
       WRITE(6,1001)
 1001 FORMAT(' This program will use the integral LET',
              ' spectrum (something.LET, generated by ',
           /,' the CREME96 codes) and device'
     &
              ' characteristics (input below) to calculate',
            /,' a heavy-ion induced SEU rate (in upsets/bit'
             ' /sec or /day).',
            /,' Before running this program you must do:',
                        to generate the environment',
             FLUX
             ' outside of the spacecraft;',
     & /,' TRANS to transport the particle fluxes',
           ' through shielding; and',
     & /,' LETSPEC to create an integral LET spectrum.'
         /,' For many devices and applications you should also do:',
             PUP to calculate the rate',
            ' of proton-induced SEUs.',
     &//,' NOTE: Before running this or any other CREME96 programs',
     & ' please define three ',
     & /,' logicals: ',/,
     & /,4x,' CREME96
                            as the directory where CREME96 source',
              ' & executables reside.',
     & /,4x,' CR96TABLES as the directory in which CREME96 data',
              ' tables reside.',
     & /,4x,' USER
                            as the directory in which output files',
              ' should be written.',
     & //,' Now begin specification of inputs for the',
```

```
LET FILE='
           CONTINUE
     110
           CALL RETRY_INPUT(IERR)
           WRITE (6, 1100)
           FORMAT(' Enter name of integral LET spectrum file',
    1100
            ' (something.LET):')
           READ(*,1,ERR=110,IOSTAT=IERR) LET FILE
           FORMAT (A80)
    1
          IF (LET FILE.EQ. IBLANK) THEN
              WRITE (6, 1914)
              FORMAT(1x,' You must specify an input .LET file',
    1914
                        ' from LETSPEC DRIVER here. ',
        &
                         ' Please try again.',/)
        &
               CALL CHECK FILE (IFILETYPE, LET FILE, IACCEPT)
               GOTO 110
          ELSE
              IFILETYPE=5
              WRITE(6,1110) LET FILE
              FORMAT (' Input LET File = ',/,1x,A80)
    1110
              CALL CHECK FILE (IFILETYPE, LET FILE, IACCEPT)
              IF (IACCEPT.NE.0) GOTO 110
ENDIF
     120 CONTINUE
         CALL RETRY INPUT (IERR)
         WRITE(6,1120)
1120 FORMAT(1x,' Enter name for an output file, which will',
& ' record the inputs and results. ',
53
        & /,1x,' (If no report file is wanted, hit <CARRIAGE RETURN>.)')
14
     121 CONTINUE
READ(*,1,ERR=120,IOSTAT=IERR) REPORT FILE
Lj
į.,.....
          IF (REPORT_FILE(1:2).EQ.'-1') GOTO 110
ű
          IF (REPORT FILE.EQ.IBLANK) THEN
              REPORT FILE='NULLFILE'
              WRITE (6, 1121)
    1121
              FORMAT(1x,' No report file created by HI UPSET DRIVER.')
          ELSE
              CALL CHECK OUTPUT FILE (REPORT FILE, IACCEPT)
              IF (IACCEPT.NE.0) GOTO 121
              WRITE(6,1122) REPORT FILE(1:79)
              FORMAT(1x,' Report file created by HI_UPSET_DRIVER: ',
    1122
                   /,1x,A79)
          ENDIF
          ENDIF
          CALL GET_HI_XS_INPUTS(DEVICE_LABEL,XM,YM,ZM,FUNNEL,NBITS,
                                IPARAM, PARAMS, XSECT FILE)
          IF (IENTER.EQ.1 .and. DEVICE LABEL(1:2).EQ.'-1') GOTO 120
          RETURN
```

' SEU rate calculation: ')

```
SUBROUTINE INITIALIZE PROTON UPSETS (PROTON FILE, NBITS,
     æ
                                           IPARAM, PARAMS, XSECT FILE,
     æ
                                           IENTER,
     æ
                                           DEVICE LABEL, REPORT FILE)
C
C
C
        Generates interactive dialogue to get necessary input parameters
С
        for proton-induced SEU rate:
C
C
       Written by:
                      Allan J. Tylka
C
                      Code 7654
C
                      Naval Research Laboratory
C
                      Washington, DC 20375-5352
C
                      tylka@crs2.nrl.navy.mil
C
C
       Last update: 20 August 1996
C
C-----
C
       IMPLICIT NONE
       CHARACTER*80 PROTON FILE, XSECT FILE, REPORT FILE
       CHARACTER*40 DEVICE LABEL
       REAL*4 PARAMS, NBITS
       INTEGER*4 IPARAM, IENTER, IFILETYPE, IACCEPT
       DIMENSION PARAMS (4)
       CHARACTER*1 IBLANK
       DATA IBLANK/' '/
       INTEGER*4 IERR
       DATA IERR/0/
      IF (IENTER.EQ.1) THEN
      WRITE(6,1000)
1000 FORMAT (' CREME96 IONIZING RADIATION ENVIRONMENT MODEL',
    & /,' --> Proton-Induced Single Event Upset'.
    & ' (SEU) Rate Calculation')
      WRITE (6,1001)
1001 FORMAT(' This program will use the differential',
             ' proton flux generated by CREME96',
           /,' [something.TFX or (for zero shielding) something.FLX',
    &
             ' or .tr*]',
    &
            ' and device characteristics',
           /,' [input below] to calculate a proton-induced',
            ' SEU rate (in SEUs/bit/sec or /day).',
    æ
           /,' NOTE: the .TFX/.FLX file may contain other species',
            ' in addition to protons, but',
           /,' they will be ignored here.',
    æ
          //,' Before running this program, you must do:',
    æ
           /,' FLUX to generate the particle',
    æ
            ' environment outside of the spacecraft;',
    &
           /,' TRANS to transport the',
    Æ
            ' particle fluxes through shielding.',
    £
           /,' For many devices and applications',
            ' you should also do: ',
           /,' LETSPEC and HUP to calculate the rate of',
           ' heavy-ion-induced SEUs.',
    &//,' NOTE: Before running this or any other CREME96 programs',
    & ' please define three ',
    & /,' logicals: ',/,
    & /,4x,' CREME96
                        as the directory where CREME96 source',
```

```
& /,4x,' CR96TABLES as the directory in which CREME96 data',
                 ' tables reside.',
        & /,4x,' USER
                               as the directory in which output files',
                 ' should be written.',
        & //,' Now begin specification of inputs for the',
              ' proton-SEU rate calculation: ')
           CONTINUE
           CALL RETRY INPUT (IERR)
           WRITE(6,1100)
           FORMAT(' Enter name of flux file containing the proton spectrum',
    1100
                  ' (something.TFX or .FLX):')
           READ(*,1,ERR=110,IOSTAT=IERR) PROTON FILE
           FORMAT (A80)
    1
           IF (PROTON_FILE.EQ.IBLANK) THEN
              WRITE(6,1914)
              FORMAT(1x,' You must specify either an input .TFX file',
    1914
                        ' from TRANSPORT_DRIVER or',
        &
              /,lx,' (in the case of zero shielding) an input',
        &
                        ' .FLX file from FLUX DRIVER',
              /,1x,' Please try again.',/)
               GOTO 110
         ELSE
IFILETYPE=4
              WRITE(6,1110) PROTON_FILE
              FORMAT(' Input Particle Flux File (containing protons) = '
    1110
                     ,/,1x,A80)
              CALL CHECK FILE (IFILETYPE, PROTON FILE, IACCEPT)
              IF (IACCEPT.NE.0) GOTO 110
          ENDIF
į.
*
    120 CONTINUE
į.
          CALL RETRY INPUT (IERR)
         WRITE(6,1120)
ij.
    1120 FORMAT(1x,' Enter name for an output file, which will',
        & ' record the inputs and results. ',
        & /,1x,' (If no report file is wanted, hit <CARRIAGE RETURN>.)')
     121 CONTINUE
          READ(*,1,ERR=120,IOSTAT=IERR) REPORT FILE
          IF (REPORT FILE(1:2).EQ.'-1') GOTO 110
          IF (REPORT FILE.EQ. IBLANK) THEN
              REPORT FILE='NULLFILE'
              WRITE (6, 1121)
    1121
              FORMAT(1x,' No report file created by HI UPSET DRIVER.')
          ELSE
              CALL CHECK_OUTPUT_FILE (REPORT_FILE, IACCEPT)
              IF (IACCEPT.NE.0) GOTO 121
              WRITE(6,1122) REPORT FILE(1:79)
    1122
              FORMAT(1x,' Report file created by HI UPSET DRIVER: ',
                   /,1x,A79)
          ENDIF
          ENDIF
```

CALL GET PROTON XS INPUTS (DEVICE LABEL,

' & executables reside.',

NBITS, IPARAM, PARAMS, XSECT_FILE)

IF (IENTER.EQ.1 .and. DEVICE_LABEL(1:2).EQ.'-1') GOTO 120

RETURN END

```
XM, YM, ZM, FUNNELM,
     &
                                                SEU RATE)
     æ
C
        Subroutine for performing numerical integration over the soft-turn
C
        on in the heavy-ion LET-dependent cross-section
С
C
                        NPTS = number of datapoints in input array
C
        INPUTS:
                        LETG = array containing LET values in MeV-cm2/g
С
                        FLUX = array containing the flux (in /m2-s-sr)
С
                                 of particles with LET > LETG
C
                        XSECT = array of cross-section values (in 1.0E-8 cm2/bit)
C
                                 corresponding to the LETG values
C
                        XM, YM, ZM = device dimensions in microns
C
                                    (ZM = depth of sensitive area,
C
                                          typically 0.5-2.0 microns)
C
                        FUNNELM = funnel length (in microns)
С
C
                        SEU RATE = SEU rate in SEUS/s/bit
C
        OUTPUT:
C
                        Allan J. Tylka
C
        Written by:
                        Code 7654
C
                        Naval Research Laboratory
C
                        Washington, DC 20375-5352
C
С
                        tylka@crs2.nrl.navy.mil
C
С
                       24 October 1996
        Last update:
C
C--
        IMPLICIT NONE
        INTEGER*4 NPTS, NPTSMAX, K
        PARAMETER (NPTSMAX=5000)
        REAL*4 LETG, FLUX, XSECT, XM, YM, ZM, FUNNELM, SEU RATE, QC, AFRAC,
                UPS1, UPS2, DELTA U, DELTA_U_SAVE, AFRACMAX
        DATA AFRACMAX/0.99/
        DIMENSION LETG(1), FLUX(1), XSECT(1)
        DIMENSION QC (NPTSMAX), AFRAC (NPTSMAX)
         INTEGER*4 NERRORS, QPTS, KLAST
        LOGICAL QUIET
        DATA QUIET/.true./
C
C
         Integrates over the soft turn-on in the heavy-ion SEU cross-section
         by calculating rate at each critical charge value and adding
С
         together in a sum weighted by the corresponding cross-section
C
C
         CALL SAMPLE SOFT TURN ON (NPTS, NPTSMAX,
                                   LETG, XSECT, XM, YM, ZM, FUNNELM,
     &
                                   AFRACMAX,
     &
                                   QPTS, QC, AFRAC)
     æ
C
         Now calculate upsets for each critical charge interval
C
         NERRORS=0
         SEU RATE=0.0
         CALL GET UPSET (XM, YM, ZM, FUNNELM, QC(1), NPTS, LETG, FLUX, UPS1)
         DO 200 K=1,QPTS-1
```

SUBROUTINE INTEGRATE_HEAVY_ION_UPSETS(NPTS, LETG, FLUX, XSECT,

```
The state of the s
```

```
IF (DELTA U.LT.O.) THEN
С
           10/24/96: The original versions of CREME and CREME96 contained
С
C
           an occasional bug here. The SEU rate should, of course,
C
           be a monotonically decreasing function of increasing critical
           charge. Due to poor sampling around LETs corresponding to the peaks
С
С
           in the differential pathlength distribution, occasionally
           the rates returned from GET UPSET did not show this. This problem
С
С
           has now (I believe) been fixed, by explicitly installing a
           higher density sampling around these values. The following
C
C
           error message should therefore NEVER be activated. AJT
           NERRORS=NERRORS+1
           IF (.not. quiet) WRITE(6,9999) DELTA_U,K, QC(K), UPS1,
                                           K+1,QC(K+1),UPS2,
     &
                                           AFRAC(K), AFRAC(K+1)
     æ
 9999
                        ERROR in SEU vs. QC: DELTA U = ',E13.6,
           FORMAT(1x,'
                /,1x,'
                        K ,QC(K) ,UPS1: ', I6,1x,E13.6,5x,E13.6,
     &
                /,1x,' K+1,QC(K+1),UPS2: ', I6,1x,E13.6,5x,E13.6,
                /,lx,' AFRAC(K),AFRAC(K+1): ',E13.6, 5x,E13.6,
                /,1x,' PLEASE NOTIFY tylka@crs2.nrl.navy.mil.')
           ELSE
           DELTA U SAVE=DELTA U
           ENDIF
           Protect against bug of undetermined origin:
С
           IF (DELTA U.LT.O.) DELTA U=DELTA U SAVE
           SEU RATE=SEU RATE+DELTA U
           IF (AFRAC(K+1).GE.AFRACMAX) THEN
               Plateau in cross-section reached. Terminate numerical
               integration over soft turn-on.
               KLAST=K+1
               GOTO 210
           ENDIF
C
           Store SEU rate for next integration step:
           UPS1=UPS2
 200
        CONTINUE
        KLAST=QPTS
        CONTINUE
 210
        CALL GET UPSET (XM, YM, ZM, FUNNELM, QC (KLAST), NPTS, LETG, FLUX, UPS1)
        IF (UPS1.LT.0.) UPS1=0.0
        SEU RATE=SEU RATE+UPS1*AFRAC(KLAST)
C
        IF (.not.quiet.and.NERRORS.GT.0.)WRITE(6,9990) QPTS,NERRORS
        FORMAT(1X, 'Debug I H I U: NPTS, NERRORS = ',215)
 9990
        RETURN
        END
```

CALL GET UPSET (XM, YM, ZM, FUNNELM, QC (K+1), NPTS, LETG, FLUX, UPS2)

DELTA U= (UPS1-UPS2) *0.5* (AFRAC(K) +AFRAC(K+1))

```
SUBROUTINE SAMPLE SOFT TURN ON (NPTS, NPTSMAX,
     &
                                         LETG, XSECT, XM, YM, ZM, FUNNELM.
     &
                                         AFRACMAX,
     &
                                         QPTS, QC, AFRAC)
         IMPLICIT NONE
        INTEGER*4 NPTS, NPTSMAX, QPTS
        REAL*4 LETG, XSECT, XM, YM, ZM, FUNNELM, AFRACMAX, QC, AFRAC
        DIMENSION LETG(1), XSECT(1), QC(1), AFRAC(1)
        INTEGER*4 K, NSAMPMAX, KMOD, KLASTO, KFIRST, KLAST
        REAL*4 AFRACTEST
        DATA NSAMPMAX/100/
        Comparison of SEU rates for various devices showed that sampling
C
        the turn on region of the cross-section curve in 100-200 points
C
C
        should be adequate. NSAMPMAX is used to reduce the number of
C
        points in the cross-section sampling to this level.
C
        OPTS=1
        DO 50 K=2, NPTS
           IF (XSECT(K).GT.O.) THEN
                QPTS=QPTS+1
               KLAST0=K
                AFRACTEST=XSECT(K)/(XM*YM)
                IF (AFRACTEST.GE.AFRACMAX) GOTO 51
           ENDIF
 50
        CONTINUE
        CONTINUE
        KMOD=QPTS/NSAMPMAX
        IF (KMOD.LE.1) KMOD=1
        QPTS=1
        KFIRST=1
        DO 100 K=2, NPTS, KMOD
           IF (XSECT(K).GT.O.) THEN
               IF (QPTS.EQ.1) KFIRST=K-1
               Now convert LET values in cross-section table to critical
C
С
               charges, in picocoulombs. See SEE Notebook #1, p. 5
               QPTS=QPTS+1
               QC(QPTS) = LETG(K) * (ZM+FUNNELM)*1.033E-5
               Also scale cross-section value by nominal area:
C
               AFRAC (QPTS) = XSECT (K) / (XM*YM)
               KLAST=K
               IF (AFRAC (QPTS).GE.AFRACMAX) GOTO 101
               IF (QPTS.EQ.NPTSMAX) THEN
                WRITE(6,9998) QPTS
 9998
                FORMAT(1x,' CAUTION from INTEGRATE_HEAVY_ION_UPSETS:',
     &
                         /,'
                                Maximum Array size reached: ',
                                QPTS = ', I4)
                GOTO 101
               ENDIF
           ENDIF
100
        CONTINUE
101
        CONTINUE
        Make sure we catch the plateau:
        IF (KLAST.LT.KLASTO) THEN
            IF (QPTS.LT.NPTSMAX) THEN
              QPTS=QPTS+1
```

```
AFRAC (QPTS) =XSECT (KLASTO) / (XM*YM)
              QC(QPTS) = LETG(KLASTO) * (ZM+FUNNELM) *1.033E-5
            ENDIF
        ELSE
            KLAST0=KLAST
        ENDIF
        Also store last value with zero cross-section:
C
        QC(1)=LETG(KFIRST)*(ZM+FUNNELM)*1.033E-5
        Following should be zero:
С
        AFRAC(1)=XSECT(KFIRST)/(XM*YM)
        RETURN
```

```
SUBROUTINE INTEGRATE_PROTON_UPSETS(NPTS,EN,FLUX,XSECT,SEU_RATE)
C
      Performs numerical integration of flux X cross-section integral
C
       for proton-induced SEUs.
С
С
       Inputs: NPTS = number of points in input arrays
С
               EN = array of proton energies (in MeV)
С
               FLUX = array of proton differential flux
C
                      (in protons/m2-s-sr-MeV), evaluated at EN
С
               XSECT= array of SEU cross-sections, in 1.0E-12 cm2/bit,
С
                      evaluated at EN
С
       Output: SEU_RATE = #SEUs/s/bit
С
C
С
                      Allan J. Tylka
        Written by:
С
                      Code 7654
C
                      Naval Research Laboratory
C
                      Washington, DC 20375-5352
C
                      tylka@crs2.nrl.navy.mil
C
C
        Last update: 29 March 1996
C
C
C-----
С
        IMPLICIT NONE
        INTEGER*4 I, NPTS
        REAL*4 EN, FLUX, XSECT
        REAL*4 FOURPI, DE, XINT, SEU RATE
        DIMENSION EN(1), FLUX(1), XSECT(1)
        FOURPI=16.0*ATAN(1.0)
        SEU RATE=0.0
        DO 200 I=1, NPTS-1
        DE=EN(I+1)-EN(I)
        XINT=0.5*(FLUX(I)*XSECT(I) + FLUX(I+1)*XSECT(I+1))
        SEU RATE=SEU_RATE+XINT*DE
        CONTINUE
 200
        Now apply some factors:
C
        Convert flux from /m2 to /cm2: 1.0E-4
С
        Cross-section in units of 1.0E-12 cm2
C
        Effective geometry factor = 4pi
С
        Planar detector has a geometry factor of 2pi (top & bottom);
С
        but the loss of projected area due to the obliquity factor is
С
        compensated for by the sec-theta increase in pathlength through
С
        the sensitive volume.
C
        SEU_RATE=FOURPI*SEU_RATE*1.0E-16
        RETURN
        END
```

```
REAL*4 FUNCTION INTERPOLATE XSECT TABLE (NSV, XV, YV, E)
C
С
        Function does linear interpolation in a table to evaluate
C
        the SEU cross-section
C
С
         Inputs: NSV:
                         number of entries in the table
C
                         x-coordinates of table
                  XV:
C
                  YV:
                         y-coordinates of table
C
                  E:
                         x-value at which cross-section is required
С
         Output: SEU cross-section
C
C
         NOTE: returned value will be zero at E < XV(1)
С
                returned value will be YV(NSV) at E > YV(NSV)
C
                otherwise, linearly-interpolated. It is the user's
C
                responsibility to make sure the table is ordered as
С
                monotonically-increasing XV values.
C
C
C
                       Allan J. Tylka
        Written by:
                       Code 7654
C
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С
                       tylka@crs2.nrl.navy.mil
C
        Last update:
C
                       29 March 1996
C
        IMPLICIT NONE
        INTEGER*4 NSV, I
        REAL*4 XV, YV, E
        DIMENSION XV(1), YV(1)
        INTERPOLATE_XSECT_TABLE = 0.
        IF (NSV.LE.O .OR. E.LT.XV(1)) RETURN
        IF (E.GT.XV(NSV)) THEN
            INTERPOLATE_XSECT_TABLE=YV(NSV)
        ELSE
            DO 100 I=2, NSV
               IF (E.LT.XV(I)) THEN
               INTERPOLATE XSECT TABLE=
               YV(I-1) + (E-XV(I-1)) * (YV(I)-YV(I-1)) / (XV(I)-XV(I-1))
               GOTO 200
               ENDIF
100
            CONTINUE
200
            CONTINUE
        ENDIF
        IF (INTERPOLATE_XSECT_TABLE.LT.O.) INTERPOLATE_XSECT_TABLE=0.0
        RETURN
        END
```

```
IMPLICIT NONE
      CHARACTER*80 INFILE, OUTFILE
      REAL*4 LETMINMG, LETMAXMG, LETMIN, LETMAX, ELOWER, EUPPER
      REAL*4 EMINCUT, EMAXCUT
      CHARACTER*12 TARGET
      INTEGER*4 MARR, NELM, LARR, IZMIN, IZMAX, IZLO, IZUP, M, L
      PARAMETER (MARR=5000, NELM=92, LARR=1002)
      REAL*4 INPUT FLUX (NELM, MARR), SPECT (LARR), DIFSPEC (LARR)
      INTEGER*4 VERSION_NUMBER, PROGRAM_CODE, IDIFSPEC
C
C
      Get parameters of LETSPEC calculation:
С
С
      CALL INILET(INFILE, LETMINMG, LETMAXMG, IZMIN, IZMAX, EMINCUT,
                   EMAXCUT, TARGET, OUTFILE, IDIFSPEC)
      Unload input particle flux file into array:
C
C
      CALL UNLOAD_PARTIAL_FLUX(INFILE, IZMIN, IZMAX, EMINCUT, EMAXCUT,
                                  ELOWER, EUPPER, M, IZLO, IZUP,
                                  INPUT FLUX)
С
С
      Now do integral LET spectrum calculation:
С
      CALL CREME96 LETSPEC (LETMINMG, LETMAXMG, TARGET,
                     ELOWER, EUPPER, M, IZLO, IZUP,
                     INPUT FLUX,
     &
                     VERSION NUMBER, PROGRAM CODE, IDIFSPEC,
     &
                     LETMIN, LETMAX, L, SPECT, DIFSPEC)
С
C
      Now write integral LET spectrum to output file:
C
      CALL OUTPUT_CREME96_LETSPEC(LETMIN, LETMAX, L,
                                     IZLO, IZUP, EMINCUT, EMAXCUT, TARGET,
                                     VERSION NUMBER, PROGRAM_CODE,
                                     INFILE,
                                     SPECT,
                                     OUTFILE)
С
     Also output differential LET spectrum
C
C
       IF (IDIFSPEC.EQ.1)
      *CALL OUTPUT_CREME96_DIFLET(LETMIN, LETMAX, L,
                                     IZLO, IZUP, EMINCUT, EMAXCUT, TARGET,
                                     VERSION NUMBER, PROGRAM CODE,
                                     INFILE,
                                     DIFSPEC,
                                     OUTFILE)
C
       STOP
       END
```

```
SUBROUTINE LOAD GTF (GTRANSFILE)
C
        Loads geomagnetic transmission function from specified file
C
        into a common block for later use.
C
C
        IMPLICIT NONE
        CHARACTER*80 GTRANSFILE
        INTEGER*4 NGTF, IGTF, I, IGTFUNIT, IVER, NHEADER, STAT, CREME96_OPEN
        REAL*4 R,GTF
        PARAMETER (NGTF=1001)
        COMMON/GTFDAT/IGTF, R (NGTF), GTF (NGTF)
        LOGICAL IEXIST, CREME96_INQUIRE
        DATA IGTFUNIT/15/
C
        First see if GTRANSFILE exists:
C
         INOUIRE(FILE='USER:'//GTRANSFILE, EXIST=IEXIST)
С
        iexist = creme96_inquire(gtransfile,'user')
        IF (.NOT. IEXIST) THEN
            WRITE(6,999) GTRANSFILE
            FORMAT(' Geomagnetic Transmission File = ',/,
  999
                     1x,A80,/' not found. Job aborted.')
     &
            STOP
        ENDIF
        CALL CHECK CREME96_VERSION(GTRANSFILE, IVER)
         OPEN (UNIT=IGTFUNIT, READONLY, SHARED, STATUS='OLD',
C
               FILE='USER:'//GTRANSFILE)
С
        stat = creme96_open(gtransfile,'user',igtfunit,'old')
С
        Get pass header lines:
C
        IF (IVER.EQ.101) THEN
            READ(IGTFUNIT,*)
            READ (IGTFUNIT, *)
         ELSEIF (IVER.GE.102) THEN
            READ (IGTFUNIT, *) NHEADER
            DO I=1, NHEADER
               READ (IGTFUNIT, *)
            ENDDO
         ENDIF
C
            Now begin real read-in.
            DO 5 I=1, NGTF
               READ(IGTFUNIT, *, END=6) R(I), GTF(I)
 5
            CONTINUE
            CONTINUE
 6
            IGTF=I-1
            CLOSE (15)
         RETURN
         END
```

```
SUBROUTINE LOAD SEP QSTATES
C
        Loads Solar Energetic Particle Ionic Charge State Distribution
C
        from datafile into COMMON/SEP_QSTATES
C
        11-17-97: IMPLICIT NCNE and variable-type declarations added.
С
        IMPLICIT NONE
        REAL*4 SEP QSTATES
        COMMON/SEP QSTATES/SEP_QSTATES(30,30)
        INTEGER*4 ISQUNIT
        DATA ISQUNIT/18/
        LOGICAL IEXIST, CREME96 INQUIRE
        INTEGER*4 STAT, CREME96_OPEN
        INTEGER*4 J, NLINES, ILINE, KMIN, KMAX, IDUM, K
        First see if QSTATES file is available here:
C
         INQUIRE(FILE='CREME96:QSTATE.DAT',EXIST=IEXIST)
        iexist = creme96_inquire('qstate.dat','cr96tables')
        IF (.NOT.IEXIST) THEN
             WRITE(6,999)
             FORMAT(' File CREME96:QSTATE.DAT not found. Abort.')
 999
        ELSE
         OPEN (UNIT=ISQUNIT, READONLY, SHARED, STATUS='OLD',
C
              FILE='CREME96:QSTATE.DAT')
C
        stat = creme96 open('qstate.dat','cr96tables',isqunit,'old')
        DO 500 J=1,30
           NLINES=(J-1)/8+1
           DO 400 ILINE=1, NLINES
              KMIN=(ILINE-1)*8+1
              KMAX=ILINE*8
              IF (KMAX.GT.J)KMAX=J
              READ (ISQUNIT, 9006) IDUM, (SEP QSTATES (J, K), K=KMIN, KMAX)
              FORMAT (13,8F8.5)
9006
           CONTINUE
 400
 500
        CONTINUE
        CLOSE (ISQUNIT)
        ENDIF
        RETURN
```

```
SUBROUTINE LOAD TRAPPED PROTONS (TRAPDFILE)
С
        Subroutine to unload CREME96 trapped proton spectrum from specified
C
С
        file into a common bloc, for later combination with non-trapped fluxes.
C
C
                        Allan J. Tylka
        Written by:
C
                        Code 7654
C
                        Naval Research Laboratory
С
                        Washington, DC 20375-5352
C
                        tylka@crs2.nrl.navy.mil
C
        Last update:
                        18 November 1997
C
C
С
C-
C
        IMPLICIT NONE
        CHARACTER*80 TRAPDFILE, ILINE
        INTEGER*4 MAXSPEC, N, NZ, NZT, i, ITRPSPEC
        INTEGER*4 IVER, NHEADER, STAT, CREME96 OPEN, ILONG
        REAL*4 ENTRP, FLUXTRP, EL, EU
        PARAMETER (MAXSPEC=5000)
        COMMON/TRPDAT/ITRPSPEC, ENTRP (MAXSPEC), FLUXTRP (MAXSPEC)
        CALL CHECK_CREME96_VERSION(TRAPDFILE, IVER)
        stat = creme96 open(TRAPDFILE, 'user', 10, 'old')
        ILONG=INDEX(TRAPDFILE,'.')
        IF (TRAPDFILE(ILONG+1:ILONG+2).EQ.'TR' .or.
            TRAPDFILE(ILONG+1:ILONG+3).EQ.'tr') THEN
            IF (IVER.GE.102) THEN
            READ(10,*) NHEADER
            DO i≈1,NHEADER
               READ(10,110) ILINE
 110
               FORMAT (A80)
            ENDDO
            ENDIF
            read(10,*) el,eu,n,nz,nzt
            IF (NZ.NE.1) THEN
            WRITE (6,999)
            FORMAT(1x,' WARNING: No proton spectrum in input file:',
 999
                  /,1x,' STOP in LOAD_TRAPPED_PROTONS')
            STOP
            ENDIF
C
C
            Calculate abscissae (energy values)
            ENTRP(1) = e1
            ENTRP(N)=eu
            do 100 i=2,N-1
            ENTRP(i) = el*(eu/el)**(float(i-1)/float(n-1))
100
C
C
            Read blank line
```

```
read(10,110) ILINE
            read in the flux
С
            IF (N.GT.MAXSPEC) N=MAXSPEC
            read(10,*)(fluxtrp(i),i=1,n)
            CLOSE(10)
        Eliminate end-of-file zeroes from returned spectrum:
С
        ITRPSPEC=0
        DO 1000 I=1, N
           IF (FLUXTRP(I).GT.0.0) ITRPSPEC=I
 1000
        CONTINUE
        ELSE
          WRITE(6,9999) TRAPDFILE
          FORMAT(1x,' Specified TRAPPED PROTON FILE = ',
 9999
               /,1x,A80,
               /,lx,' does not appear to be a CREME96-generated file.',
     &
               /,1x,' STOP')
        ENDIF
        RETURN
        end
```

```
REAL*4 FUNCTION MAGNETIC_RIGIDITY(EK,Q,A)
C
        Function to calculate the magnetic rigidity (in GV/c)
С
С
        given inputs:
                  EK = kinetic energy per nucleon in MeV/amu
C
C
                  Q = charge
                  A = mass number (> 0 for ions; 0 for electrons)
C
C
        IMPLICIT REAL*8 (D)
        REAL*4 EK,Q,A
        DATA DAMU/0.9315016D0/
        DATA DELEC/0.00051099906/
        DK=EK/1000.D0
        DA=A
        DQ=Q
C
        DR= gamma*beta
        IF (DA.GT.0.0) THEN
C
        Ion case:
        DR = (1.D0 + DK/DAMU) **2-1.D0
        DR=DSQRT (DR) *DA/DQ*DAMU
        ELSEIF (DA.EQ.0) THEN
        Electron case:
        DR=(1.D0+DK/DELEC)**2-1.D0
        DR=DSQRT (DR) *DELEC
        ENDIF
        MAGNETIC_RIGIDITY=DR
        RETURN
```

```
C
            Makes two-column energy vs. dE/dx tables from CREME96 software.
    C
    С
            IMPLICIT NONE
            INTEGER*4 NBINMAX, STAT, CREME96_OPEN
            PARAMETER (NBINMAX=5000)
            INTEGER*4 NBINS, K, IZ, OUTUNIT, IMAT
            REAL*4 E(NBINMAX)
            REAL*4 EMIN, EMAX, EN, DE, Z, AN, DEDX, STPOW
            CHARACTER*80 OUTFILE
            CHARACTER*12 MATERIAL, MATS(2)
            DATA MATS/'ALUMINUM','SILICON
            DATA OUTUNIT/8/
            INTEGER*4 IERR, IACCEPT
            DATA IERR/0/
            MATERIAL='ALUMINUM
            WRITE(6,8000)
            FORMAT(' This program will make a table of stopping power',
     8000
                  ' in aluminum for the specified'
         ራ
                /,' nuclei. The table will be specified for NBINS',
                  ' logarithmically-spaced energy',
                /,' values between limits EMIN and EMAX (in MeV/nuc)')
            CONTINUE
     8101
            CALL RETRY INPUT (IERR)
THE SHARE STATE
            WRITE(6,8001) NBINMAX
            FORMAT(' Enter EMIN, EMAX (in MeV/nuc) and NBINS (<', I5,')')
     8001
            READ(*,*,ERR=8101,IOSTAT=IERR) EMIN,EMAX,NBINS
            IF (NBINS.GT.NBINMAX) NBINS=NBINMAX
            WRITE(6,8002) EMIN, EMAX, NBINS
     8002 FORMAT(1x,' EMIN = ',E13.5,' EMAX = ',E13.5,' NBINS = ',I5)
la la
=
           CONTINUE
     8103
            CALL RETRY INPUT (IERR)
П
            WRITE(6,8003)
     8003 FORMAT(' Select material: Enter 1 for Aluminum; 2 for Silicon')
            READ(*,*,ERR=8103,IOSTAT=IERR) IMAT
            MATERIAL=MATS (IMAT)
            WRITE(6,8004) MATERIAL
            FORMAT(1x,' Material = ',A12)
     8004
    C
          Compute energies on logaritmically-spaced grid
            DE= (EMAX/EMIN) ** (1./(NBINS-1.))
            E(1) = EMIN
            DO K=2, NBINS-1
                E(K) = E(K-1) *DE
             END DO
            E(NBINS) = EMAX
      9102
            CONTINUE
            CALL RETRY INPUT (IERR)
            WRITE(6,9002)
            FORMAT(1x,' Enter name of output file:')
      9002
             READ(*,2,ERR=9102,IOSTAT=IERR) OUTFILE
             FORMAT (A80)
             CALL CHECK OUTPUT_FILE (OUTFILE, IACCEPT)
             IF (IACCEPT.NE.O) THEN
```

WRITE(6,1011) OUTFILE(1:75)

program MAKE_DEDX_TABLE

```
191
            CONTINUE
     114
            CALL RETRY_INPUT(IERR)
            WRITE(6,9003)
     9003
         &
                  /,8x,' A = mass number')
     115
            CONTINUE
            Z=IZ*1.0
             WRITE(6,9004) IZ
     9004
GOTO 114
            ENDIF
            FORMAT(1x,'%Energy (MeV/nuc)'
     190
                 ' dE/dx (MeV/(g/cm2))',
last.
æ
C
            DO 80 K=1, NBINS
EN=E(K)
                DEDX=STPOW(EN, Z, AN, MATERIAL)
    С
                DEDX=AN*DEDX
                write (OUTUNIT, 34) EN, DEDX
                FORMAT (2X, E12.5, 2X, E12.5)
    34
    80
            continue
            CONTINUE
     9106
             CALL RETRY INPUT (IERR)
            WRITE(6,9006)
```

```
WRITE(6,1012)
        FORMAT(1x,' Previous try at OUTPUT name = ',/,5x,A75)
1011
        FORMAT(1x,' Try again.')
1012
        GOTO 9102
        ENDIF
        open(unit=OUTUNIT, status='new', file='USER:'//OUTFILE)
С
        stat = creme96 open(outfile, 'user', outunit, 'new')
        WRITE (OUTUNIT, 191) EMIN, EMAX, NBINS
        FORMAT(1x, '%EMIN = ', E13.5,' EMAX = ', E13.5,' NBINS = ', I5)
        FORMAT(1x,' Enter first desired IZ, A values for table: '
             /,8x,' IZ = atomic number',
        READ(*, *, ERR=114, IOSTAT=IERR) IZ, AN
        IF (IZ.LT.0 .or. IZ.GT.92) THEN
         FORMAT(1x,' Invalid atomic number: ', I6,' Please try again.')
        WRITE (OUTUNIT, 190) IZ, AN, MATERIAL
             ' for Z = ',I2,' A = ',F6.2,' in ',A12)
           STPOW calculates energy loss in MeV/nuc:
 9006
        FORMAT(1x,' Enter next IZ, A value (0 0 to end program):')
        READ(*, *, ERR=9106, IOSTAT=IERR) IZ, AN
        IF (IZ.GT.0) GOTO 115
        CLOSE (OUTUNIT)
        STOP
        end
```

```
SUBROUTINE MAKE DIFLET SPECTRUM (LETMIN, LETMAX, LIN, SPECT, DIFSPEC)
   С
          Makes SIMPLE (ie., needs work) numerical differentiation of
   C
          integral LET spectrum to produce a differential LET spectrum.
   C
   C
          IMPLICIT NONE
          REAL*4 LETMIN, LETMAX
          INTEGER*4 LIN, L, K, LARR
          REAL*4 SPECT, DIFSPEC
          DIMENSION SPECT(1), DIFSPEC(1)
          REAL*8 DY, DL, LETG
          PARAMETER (LARR=1002)
         DIMENSION LETG(LARR)
         Fill array of LET values:
   С
          L=LIN
          IF (L.GT.LARR) L=LARR
          DL=(LETMAX/LETMIN)**(1./FLOAT(L-1))
          LETG(1)=LETMIN
          DIFSPEC(1) = 0.0
          DO 400 K=2,L-1
             LETG(K)=LETG(K-1)*DL
             DIFSPEC(K) = 0.0
    400 CONTINUE
          LETG(L)=LETMAX
          DIFSPEC(L)=0.0
         Now calculate differential LET spectrum.
DL=LETG(2)-LETG(1)
          IF (SPECT(2).GT.0.)
Englis
              DIFSPEC(1) = -SPECT(1) *ALOG(SPECT(2)/SPECT(1))/DL
S
         DO 500 K=2,L-1
             IF (SPECT(K+1).GT.0.0) THEN
                 DL = (LETG(K+1) - LETG(K-1))
                 DY=-ALOG (SPECT (K+1) /SPECT (K-1))
į.i.
             ELSE
ũ
                 IF (SPECT(K).GT.0.0) THEN
                 DL=LETG(K)-LETG(K-1)
                 DY=-ALOG(SPECT(K)/SPECT(K-1))
                 ENDIF
             ENDIF
             DIFSPEC(K) = SPECT(K) *DY/DL
     500 CONTINUE
          RETURN
          END
```

```
program MAKE FLUX FIG
С
        Rewrites flux output file from CREME96/UPROP format
C
        into a .FIG file.
C
        IMPLICIT NONE
        INTEGER*4 MARR, NELM, STAT, CREME96_OPEN
        PARAMETER (MARR=5000, NELM=92)
        REAL*4 FLUX (NELM, MARR), E (MARR)
        REAL*4 EL, EU, ETEMP, EMINCUT, EMAXCUT
        INTEGER*4 IACCEPT, IFILETYPE, IZMIN, IZMAX, M, IZLO, IZUP
        CHARACTER*80 INFILE, OUTFILE
        CHARACTER*80 ILINE, TEMPLINE
        INTEGER*4 IZTARG, NHMAX, LINEMAX, ICOUNT
        PARAMETER (NHMAX=30)
        DIMENSION ILINE (NHMAX)
        INTEGER*4 I, OUTUNIT, FHDUNIT, NHEADER, IZDUM, ILONG
        LOGICAL ZERO
        INTEGER*4 IERR
        DATA IERR/0/
        OUTUNIT=42
        FHDUNIT=43
 112
        CONTINUE
        CALL RETRY_INPUT(IERR)
        WRITE (6, 9001)
        FORMAT(1x,' Enter name of input file',
 9001
                   ' (something.FLX, .TFX, or .TR*):')
        READ(*,2,ERR=112,IOSTAT=IERR) INFILE
        FORMAT (A80)
        IFILETYPE=3
        WRITE(6,1020) INFILE
 1020 FORMAT(' Input Flux File =',/,1x,A80)
        CALL CHECK_FILE (IFILETYPE, INFILE, IACCEPT)
         IF (IACCEPT.NE.0) GOTO 112
C
         Now unload fluxes from this file:
C
         EMINCUT=0.0
         EMAXCUT=1.0E+24
         CALL UNLOAD_PARTIAL_FLUX(INFILE, IZMIN, IZMAX, EMINCUT, EMAXCUT,
                                           EL, EU, M, IZLO, IZUP,
      &
                                           FLUX)
         Calculate abscissae (energies)
C
         e(1) = EL
         e(M) = EU
         do 10 i=2, M-1
            e(i) = el*(eu/el) **(float(i-1)/float(M-1))
         continue
 10
         Now start to copy information to new file:
 C
         CONTINUE
  9102
         CALL RETRY_INPUT(IERR)
         WRITE(6,9002)
         FORMAT(1x,' Enter name of output file (something.FIG):')
  9002
         READ(*,2,ERR=9102,IOSTAT=IERR) OUTFILE
```

CALL CHECK_OUTPUT_FILE (OUTFILE, IACCEPT)

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IF (IACCEPT.NE.0) THEN
        WRITE(6,1010) INFILE(1:75)
        WRITE(6,1011) OUTFILE(1:75)
        WRITE(6,1012)
        FORMAT(1x,' INPUT file = ', /, 5x, A75)
 1010
        FORMAT(1x,' Previous try at OUTPUT name = ',/,5x,A75)
 1011
        FORMAT(1x,' Try again.')
 1012
        GOTO 9102
        ENDIF
        open(unit=OUTUNIT, status='new', file='USER:'//OUTFILE)
C
        stat = creme96_open(outfile,'user',outunit,'new')
        WRITE (OUTUNIT, 185) INFILE (1:70)
        FORMAT ('*', A50)
 185
С
        Now add FIGGEN header information:
C
C
        Write FIGGEN header
         OPEN (UNIT=FHDUNIT, status='old', readonly, shared,
C
              file='CREME96:FLX FIG.header')
C
        stat = creme96 open('flx fig.header','cr96tables',fhdunit,'old')
        CONTINUE
6
        READ (FHDUNIT,2,END=8) ILINE(1)
        WRITE (OUTUNIT, 2) ILINE (1)
        GOTO 6
8
        CONTINUE
        CLOSE (FHDUNIT)
        ILONG=LEN(INFILE)
        CALL CAPITALIZE STRING (INFILE, ILONG)
        WRITE (OUTUNIT, 186) INFILE (1:50)
        FORMAT('ST 0.3 100 1.0E+5 0.//^^',A50)
 186
        Transfer header information from input file
        CALL UNLOAD HEADERS (INFILE, NHMAX, ILINE, LINEMAX)
        IF (LINEMAX.GT.0) THEN
        DO 100 I=1, LINEMAX
           TEMPLINE=ILINE(I)
           TEMPLINE='*'//TEMPLINE(2:80)
           WRITE (OUTUNIT, 2) TEMPLINE
 100
        CONTINUE
        ENDIF
 114
        CONTINUE
        CALL RETRY INPUT (IERR)
        WRITE(6,9003)
        FORMAT(1x,' Enter desired IZMIN, IZMAX for FIGure: ')
 9003
        READ (*, *, ERR=114, IOSTAT=IERR) IZMIN, IZMAX
 115
        CONTINUE
        DO 200 IZTARG=IZMIN, IZMAX
        IF (IZTARG.LT.0 .or. IZTARG.GT.92) THEN
            WRITE(6,9004) IZTARG
            FORMAT(1x,' Invalid atomic number: ', I6)
 9004
            GOTO 200
        ENDIF
C
        Does this file contain the Z value of interest?
C
        IF (IZTARG.LT.IZLO .or. IZTARG.GT.IZUP) THEN
           WRITE (6,9005) IZLO, IZUP, IZTARG
```

```
/,1x, 'IZ = ',I5,' not found here.')
                GOTO 200
             ENDIF
             WRITE (OUTUNIT, 30) IZTARG
             FORMAT('*IZ = ',I3)
     30
             ZERO=.TRUE.
             DO 50 I=1, M
                IF (FLUX(IZTARG, I).GT.0.) THEN
                    ZERO=.FALSE.
                    GOTO 55
                ENDIF
             CONTINUE
     50
     55
             CONTINUE
             IF (.NOT.ZERO) THEN
             IF (MOD(IZTARG,2) .EQ.0) THEN
                     WRITE (42,31) M
             ELSE
                     WRITE(42,32) M
             ENDIF
     31
            FORMAT ('FRENCH', 14,' 0.01 0')
            FORMAT ('FRENCH', 14,' 0.10 2')
     32
            DO 80 I=1,M
             IF (FLUX(IZTARG, I).GT.0.0) THEN
                 write(OUTUNIT,34) e(i),flux(IZTARG,i)
             ENDIF
             FORMAT (2X, E12.5, 2X, E12.5)
    34
             continue
    80
35
             ICOUNT=ICOUNT+1
             IF (ICOUNT.GT.3) ICOUNT=1
ETEMP=E (M) * (1.+ICOUNT*0.05)
L.
             WRITE (OUTUNIT, 90) ETEMP, FLUX (IZTARG, M), IZTARG
    90
             FORMAT ('STHC -0.2 ', E12.5, 2x, E12.5, ' 0//', I2)
             ENDIF
    200
             CONTINUE
             CLOSE (OUTUNIT)
             STOP
             end
```

9005

FORMAT(1x,' In this file: IZLO = ', I5,' IZHI = ', I5,

```
program MAKE_FLUX_TABLE
    C
    C
            Rewrites flux output file from CREME96/UPROP format
   C
            into a two column table.
    C
            IMPLICIT NONE
            INTEGER*4 MARR, NELM, STAT, CREME96 OPEN
            PARAMETER (MARR=5000, NELM=92)
            REAL*4 FLUX (NELM, MARR), E (MARR)
            REAL*4 EL, EU, EMINCUT, EMAXCUT
            INTEGER*4 IACCEPT, IFILETYPE, IZTARG, M, IZLO, IZUP
            CHARACTER*80 INFILE, OUTFILE
            INTEGER*4 I, OUTUNIT, NHEADER, IZDUM
            INTEGER*4 IERR
            DATA IERR/0/
            OUTUNIT=42
            CONTINUE
    112
            CALL RETRY INPUT (IERR)
            WRITE (6,9001)
            FORMAT(1x,' Enter name of input file',
    9001
                  ' (something.FLX, .TFX, or .TR*):')
            READ (*, 2, ERR=112, IOSTAT=IERR) INFILE
            FORMAT (A80)
            IFILETYPE=3
            WRITE(6,1020) INFILE
    1020
            FORMAT(' Input Flux File =',/,1x,A80)
            CALL CHECK FILE (IFILETYPE, INFILE, IACCEPT)
            IF (IACCEPT.NE.0) GOTO 112
   C
   C
            Now unload fluxes from this file:
            IZDUM=0
-
            EMINCUT=0.0
15
            EMAXCUT=1.0E+24
CALL UNLOAD PARTIAL FLUX (INFILE, IZDUM, IZDUM, EMINCUT, EMAXCUT,
δe
                                              EL, EU, M, IZLO, IZUP,
FLUX)
C
            Calculate abscissae (energies)
            e(1) = EL
            e(M) = EU
            do 10 i=2, M-1
               e(i) = el*(eu/el)**(float(i-1)/float(M-1))
   10
            continue
   С
            Now start to copy information to new file:
    9102
            CONTINUE
            CALL RETRY_INPUT(IERR)
            WRITE(6,9002)
    9002
            FORMAT(1x,' Enter name of output file (something.DAT):')
            READ(*,2,ERR=9102,IOSTAT=IERR) OUTFILE
            CALL CHECK OUTPUT FILE (OUTFILE, IACCEPT)
            IF (IACCEPT.NE.0) THEN
            WRITE(6,1010) INFILE(1:75)
            WRITE(6,1011) OUTFILE(1:75)
            WRITE(6,1012)
    1010
            FORMAT(1x,' INPUT file = ',/,5x,A75)
    1011
            FORMAT(1x,' Previous try at OUTPUT name = ',/,5x,A75)
```

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1012
        FORMAT(1x,' Try again.')
        GOTO 9102
        ENDIF
        open(unit=OUTUNIT, status='new', file='USER:'//OUTFILE)
C
        stat = creme96 open(outfile,'user',outunit,'new')
        WRITE (OUTUNIT, 185) INFILE (1:79)
        FORMAT(1x,A79)
 185
        CALL CHECK_HEADER_LENGTH(INFILE, NHEADER)
        CALL COPY HEADERS (INFILE, NHEADER, OUTUNIT)
 114
        CONTINUE
        CALL RETRY INPUT (IERR)
        WRITE(6,9003)
 9003
        FORMAT(1x,' Enter desired IZ value for table: ')
        READ(*,*,ERR=114,IOSTAT=IERR) IZTARG
 115
        CONTINUE
        IF (IZTARG.LT.0 .or. IZTARG.GT.92) THEN
         WRITE(6,9004) IZTARG
 9004
         FORMAT(1x,' Invalid atomic number: ', I6,' Please try again.')
         GOTO 114
        ENDIF
C
С
        Does this file contain the Z value of interest?
        IF (IZTARG.LT.IZLO .or. IZTARG.GT.IZUP) THEN
           WRITE(6,9005) IZLO, IZUP, IZTARG
 9005
           FORMAT(1x,' In this file: IZLO = ', I5,' IZHI =', I5,
                /,1x,' IZ = ', I5,' not found here.',
     &
     &
                /,1x,' Please try again.')
           GOTO 114
        ENDIF
        WRITE (OUTUNIT, 190) IZTARG
190
        FORMAT(1x, '% Energy (in MeV/nuc)'
             ' vs. Flux (in particles/m2-s-sr-MeV/nuc)',
     &
             ' for Z = ', I2)
     &
C
        DO 80 I=1,M
        IF (FLUX(IZTARG, I).GT.0.0) THEN
            write(OUTUNIT,34) e(i),flux(IZTARG,i)
        ENDIF
34
        FORMAT (2X, E12.5, 2X, E12.5)
80
        continue
9106
        CONTINUE
        CALL RETRY INPUT (IERR)
        WRITE(6,9006)
9006
        FORMAT(1x,' Enter next Z value (0 to end program):')
        READ(*,*,ERR=9106,IOSTAT=IERR) IZTARG
        IF (IZTARG.GT.0) GOTO 115
        CLOSE (OUTUNIT)
        STOP
        end
```

```
C
        Rewrites integral LET spectrum output file from CREME96/UPROP format
С
        to a FIGGEN file
C
С
        IMPLICIT NONE
        INTEGER*4 MARR, STAT, CREME96_OPEN
        PARAMETER (MARR=5000)
        REAL*4 FLUX (MARR), E (MARR)
        INTEGER*4 IACCEPT, IFILETYPE, M
        CHARACTER*80 INFILE, OUTFILE, HEADER
        CHARACTER*80 ILINE, TEMPLINE
        INTEGER*4 NHMAX, LINEMAX
        PARAMETER (NHMAX=30)
        DIMENSION ILINE (NHMAX)
        INTEGER*4 I, OUTUNIT, FHDUNIT, IERR, ISPECTYPE, ILONG
        DATA IERR/0/
        LOGICAL INTLET, DIFLET
        OUTUNIT=42
        FHDUNIT=43
        ILONG=0
        CONTINUE
 112
        CALL RETRY INPUT (IERR)
        WRITE(6,9001)
        FORMAT(1x,' Enter name of input file: ',
 9001
             /,5x,'(something.LET for an integral LET spectrum;',
     δc
             /,5x,' something.DLT for a differential LET spectrum)')
        READ(*,2,ERR=112,IOSTAT=IERR) INFILE
        FORMAT (A80)
        INTLET=.FALSE.
        DIFLET=.FALSE.
        ILONG=INDEX(INFILE,'.')
        IF (ILONG.EQ.0) GOTO 112
        IF (INFILE(ILONG+1:ILONG+3).EQ.'LET' .or.
            INFILE(ILONG+1:ILONG+3).EQ.'let') INTLET=.TRUE.
        IF (INFILE(ILONG+1:ILONG+3).EQ.'DLT' .or.
             INFILE(ILONG+1:ILONG+3).EQ.'dlt') DIFLET=.TRUE.
        IF (.NOT.INTLET .and. .NOT.DIFLET) THEN
 111
        CONTINUE
        CALL RETRY INPUT (IERR)
        WRITE(6,9000)
        FORMAT(1x,' LET file type not known. Please specify type:',
 9000
             /,1x,' 0=integral or 1=differential')
        READ(2, *, ERR=111, IOSTAT=IERR) ISPECTYPE
        IF (ISPECTYPE.LT.0 .or. ISPECTYPE.GT.1) GOTO 111
        IF (ISPECTYPE.EQ.0) INTLET=.TRUE.
        IF (ISPECTYPE.EQ.1) DIFLET=.TRUE.
        ENDIF
        IF (INTLET) IFILETYPE=5
        IF (DIFLET) IFILETYPE=6
        WRITE(6,1020) INFILE
        FORMAT(' Input Flux File =',/,1x,A80)
 1020
        CALL CHECK FILE (IFILETYPE, INFILE, IACCEPT)
        IF (IACCEPT.NE.0) GOTO 112
```

Now unload fluxes from this file:

program MAKE LETSPEC_FIG

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C

```
CALL UNLOAD LET SPECTRUM(INFILE, E, FLUX, M)
           Now start to copy information to new file:
   C
           CONTINUE
    9102
           CALL RETRY INPUT (IERR)
           WRITE (6, 9002)
           FORMAT(1x,' Enter name of output file (something.FIG):')
    9002
           READ(*, 2, ERR=9102, IOSTAT=IERR) OUTFILE
           CALL CHECK_OUTPUT_FILE (OUTFILE, IACCEPT)
           IF (IACCEPT.NE.O) THEN
           WRITE(6,1010) INFILE(1:75)
           WRITE(6,1011) OUTFILE(1:75)
           WRITE (6, 1012)
           FORMAT(1x,' INPUT file = ',/,5x,A75)
    1010
           FORMAT(1x,' Previous try at OUTPUT name = ', /, 5x, A75)
    1011
           FORMAT(1x,' Try again.')
    1012
           GOTO 9102
           ENDIF
           open(unit=OUTUNIT, status='new', file='USER:'//OUTFILE)
   С
            stat = creme96 open(outfile,'user',outunit,'new')
           WRITE (OUTUNIT, 185) INFILE (1:70)
           FORMAT ('*', A50)
    185
T C
           Now add FIGGEN header information:
           Write FIGGEN header
  C
            IF (INTLET) HEADER='CREME96:LETSPEC_FIG.header'
   С
            IF (DIFLET) HEADER='CREME96:DLTSPEC_FIG.header'
   C
            IF (INTLET) HEADER='LETSPEC_FIG.header'
            IF (DIFLET) HEADER='DLTSPEC FIG.header'
            OPEN(UNIT=FHDUNIT, status='old', readonly, shared, file=HEADER)
   C
            stat = creme96 open(header,'cr96tables',fhdunit,'old')
            CONTINUE
            READ (FHDUNIT, 2, END=8) ILINE(1)
            WRITE (OUTUNIT, 2) ILINE (1)
            GOTO 6
   8
            CONTINUE
            CLOSE (FHDUNIT)
            ILONG=LEN (INFILE)
            CALL CAPITALIZE_STRING(INFILE, ILONG)
            IF (INTLET) WRITE (OUTUNIT, 186) INFILE (1:50)
            IF (DIFLET) WRITE (OUTUNIT, 187) INFILE (1:50)
            FORMAT ('ST 0.3 1.0E+2 1.0E+3 0.//^^', A50)
    186
            FORMAT('ST 0.3 1.0E+3 1.0E+3 0.//^^',A50)
    187
            Transfer header information from input file
   C
            CALL UNLOAD HEADERS (INFILE, NHMAX, ILINE, LINEMAX)
            IF (LINEMAX.GT.0) THEN
            DO 100 I=1, LINEMAX
               TEMPLINE=ILINE(I)
               TEMPLINE='*'//TEMPLINE(2:80)
               WRITE (OUTUNIT, 2) TEMPLINE
            CONTINUE
    100
            ENDIF
```

IF (INTLET) WRITE (OUTUNIT, 190)

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IF (DIFLET) WRITE(OUTUNIT,191)
190
        FORMAT(1x,'*%LET (in MeV-cm2/g)'
            ' vs. Integral Flux (in particles/m2-s-sr)')
        FORMAT(1x,'*%LET (in MeV-cm2/g)'
191
     & ' vs. Differential Flux (in particles/m2-s-sr-(MeV-cm2/g))')
С
        WRITE(42,31) M
        FORMAT ('FRENCH ', I4,' 0.01 0')
31
        DO 80 I=1,M
        IF (FLUX(I).GT.0.0) THEN
            write(OUTUNIT,34) e(i),flux(i)
        ENDIF
        FORMAT (2X,E12.5,2X,E12.5)
34
80
        continue
        CLOSE (OUTUNIT)
        STOP
        end
```

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```
program MAKE_LETSPEC_TABLE
C
        Rewrites integral LET output file from CREME96/UPROP format
С
        into a two column table.
С
C
        IMPLICIT NONE
        INTEGER*4 MARR, STAT, CREME96 OPEN
        PARAMETER (MARR=5000)
        REAL*4 FLUX (MARR), E (MARR), LETMIN
        INTEGER*4 IACCEPT, IFILETYPE, M
        CHARACTER*80 INFILE, OUTFILE
        INTEGER*4 I, OUTUNIT, NHEADER, IERR, ISPECTYPE, ILONG
        DATA IERR/0/
        LOGICAL INTLET, DIFLET
        OUTUNIT=42
        ILONG=0
        CONTINUE
 112
        CALL RETRY INPUT (IERR)
        WRITE (6,9001)
       FORMAT(1x,' Enter name of input file: ',
 9001
             /,5x,'(something.LET for an integral LET spectrum;',
     &
             /,5x,' something.DLT for a differential LET spectrum)')
        READ(*,2,ERR=112,IOSTAT=IERR) INFILE
        FORMAT (A80)
        INTLET=.FALSE.
        DIFLET=.FALSE.
        ILONG=INDEX(INFILE,'.')
        IF (ILONG.EQ.0) GOTO 112
        IF (INFILE(ILONG+1:ILONG+3).EQ.'LET' .or.
             INFILE(ILONG+1:ILONG+3).EQ.'let') INTLET=.TRUE.
        IF (INFILE(ILONG+1:ILONG+3).EQ.'DLT' .or.
              INFILE(ILONG+1:ILONG+3).EQ.'dlt') DIFLET=.TRUE.
        IF (.NOT.INTLET .and. .NOT.DIFLET) THEN
        CONTINUE
 111
        CALL RETRY INPUT (IERR)
        WRITE(6,9000)
        FORMAT(1x,' LET file type not known. Please specify type:',
 9000
              /,1x,' 0=integral or 1=differential')
        READ(2,*,ERR=111,IOSTAT=IERR) ISPECTYPE
         IF (ISPECTYPE.LT.0 .or. ISPECTYPE.GT.1) GOTO 111
         IF (ISPECTYPE.EQ.0) INTLET=.TRUE.
         IF (ISPECTYPE.EQ.1) DIFLET=.TRUE.
         ENDIF
         IF (INTLET) IFILETYPE=5
         IF (DIFLET) IFILETYPE=6
         WRITE(6,1020) INFILE
         FORMAT(' Input Flux File =',/,1x,A80)
  1020
         CALL CHECK_FILE (IFILETYPE, INFILE, IACCEPT)
         IF (IACCEPT.NE.0) GOTO 112
 C
         Now unload fluxes from this file:
         CALL UNLOAD LET_SPECTRUM(INFILE, E, FLUX, M)
         Now start to copy information to new file:
 C
         CONTINUE
  9201
         CALL RETRY INPUT (IERR)
```

```
WRITE(6,9101)
        FORMAT(1x,' Enter starting LET value (in MeV-cm2/g)',
9101
                  ' to be included in output table.',
    &
                /,' (For most SEU applications, LET = 100 \text{ MeV-cm}2/g',
    &
                  ' is appropriate.)')
     &
        READ(*, *, ERR=9201, IOSTAT=IERR) LETMIN
        CONTINUE
9202
        CALL RETRY INPUT (IERR)
        WRITE(6,9002)
        FORMAT(1x,' Enter name of output file (something.DAT):')
9002
        READ(*,2,ERR=9202,IOSTAT=IERR) OUTFILE
        CALL CHECK OUTPUT FILE (OUTFILE, IACCEPT)
        IF (IACCEPT.NE.O) THEN
        WRITE(6,1010) INFILE(1:75)
        WRITE(6,1011) OUTFILE(1:75)
        WRITE(6,1012)
        FORMAT(1x,' INPUT file = ',/,5x,A75)
1010
        FORMAT(1x,' Previous try at OUTPUT name = ',/,5x,A75)
 1011
        FORMAT(1x,' Try again.')
1012
        GOTO 9202
        ENDIF
        open(unit=OUTUNIT, status='new', file='USER:'//OUTFILE)
        stat = creme96 open(outfile,'user',outunit,'new')
        WRITE (OUTUNIT, 185) INFILE (1:50), LETMIN
185
        FORMAT(1x, A50, 'LET > ', E12.5)
        CALL CHECK HEADER LENGTH (INFILE, NHEADER)
        CALL COPY HEADERS (INFILE, NHEADER, OUTUNIT)
        IF (INTLET) WRITE (OUTUNIT, 190) LETMIN
        IF (DIFLET) WRITE (OUTUNIT, 191) LETMIN
        FORMAT(1x,'%LET (in MeV-cm2/g)'
190
             ' vs. Integral Flux (in particles/m2-s-sr);',
     &
             ' LET >',E10.3)
191
        FORMAT(1x,'%LET (in MeV-cm2/g)'
     & ' vs. Diff. Flux (in #/m2-s-sr-(MeV-cm2/g));',
     & 'LET >',E10.3)
C
        DO 80 I=1,M
        IF (FLUX(I).GT.0.0.and.E(I).GE.LETMIN) THEN
            write(OUTUNIT,34) e(i),flux(i)
        FORMAT (2X, E12.5, 2X, E12.5)
34
80
        continue
        CLOSE (OUTUNIT)
        STOP
        end
```

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______
      This subroutine computes the mean free path of any nuclide
      (IZ,IA) in material NAME at energy ENERGY (MeV/N).
       It must be linked with CREME96: ZTARGET. DAT
       INCLUDE 'CREME96: ZCOMMON.CMN'
       CHARACTER*12 NAME
       REAL NA(28), IADJ(28), NASPM(28), DENS, ETAD
       INTEGER NZ(28), IGAS, NAS
       COMMON /TBLOCK/DENS, ETAD, IGAS, NAS,
                     NZ, NA, IADJ, NASPM,
    &
                     NTOTAL, AVGZ, AVGZ2, AVGA, AVGI
    &
       REAL IA, MEAN FREE_PATH
       DATA AVOGADRO/6.022045E23/ ! particles/mole
       CALL ZTARGET (NAME)
       Compute mulitiplicative factor
       FACT=1.E27*AVGA/AVOGADRO
       AVGS=0.
       DO J=1, NAS
        NNZ=NZ(J)
         (L) AN=NAA
         CALL SMASH(IZ, IA, NNZ, AAN, ENERGY, S)
         AVGS=AVGS+NASPM(J)*S
       END DO
       AVGS=AVGS/NTOTAL
       MEAN FREE PATH=AVGS/FACT
       RETURN
       END
       SUBROUTINE SMASH(IZ, IA, JZ, JA, E, CROSS_SECTION)
     ______
       Computes nucleus-nucleus total reaction cross section
C
C
       inputs:
С
             = projectile energy (MeV/nucleon)
C
         IZ = projectile charge
С
         IA = projectile mass
С
         JZ = target charge
C
С
         JA = target mass
С
       SUBROUTINE
C
       SUBROUTINE LETAW(IZ, A, ENERGY, CROSS_SECTION)
       SUBROUTINE OVERLAP (IA, JA, CROSS SECTION)
       REAL IA, JA
       Define constants
C
       PI=3.1415927
```

Zero cross section at zero energy

SUBROUTINE MFP (ENERGY, IZ, IA, NAME, MEAN FREE PATH)

```
CROSS SECTION=0.
            RETURN
          ENDIF
  С
          proton-proton reactions
          IF (IZ*IA*JZ*JA.LE.1.02) THEN
             IF (IZ.EQ.JZ) THEN
               CROSS SECTION=PTOTAL(E,1)
               CROSS SECTION=PTOTAL(E,2)
             ENDIF
             RETURN
           ENDIF
           proton or neutron projectile
   C
           IF (IZ*IA.LE.1.02) THEN
             CALL LETAW (JZ, JA, E, CROSS_SECTION)
             RETURN
           ENDIF
           proton or neutron target
Į.
           IF (JZ*JA.LE.1.02) THEN
             CALL LETAW (IZ, IA, E, CROSS_SECTION)
             RETURN
ENDIF
           general nucleus-nucleus collision
CALL OVERLAP(IA, JA, CROSS_SECTION)
Œ
           RETURN
FI.
           END
W
           FUNCTION OVERLAP (IA, JA, CROSS_SECTION)
ũ
           subroutine OVERLAP(IA, JA, CROSS_SECTION)
           Westfall mass-changing 10*pi*r**2=67.887
           CROSS SECTION=67.887*(IA**(1./3.)+JA**(1./3.)-1.12)**2
           RETURN
           END
           FUNCTION PTOTAL (E, N)
           This function computes the proton-proton and proton-neutron
   С
           total cross section according to empirical formulas.
   C
           It is valid for energies between 40 MeV and 1000 GeV.
           If N=1 (pp), if N=2 (pn). Energy in MeV.
           DIMENSION P(12,2),C(3)
           DATA P/293.3,1.99,35.0,15.02,2.925,548.3,
        & 1104.,-.1444,4091.,.1174,75100.,.05061,
        & 1623.,1.423,30.97,13.08,.9946,561.9,
        & 2677.,-.0586,25950.,.0691,1.E6,0./
           A=(P(1,N)/E)**P(2,N) + P(3,N)
           B=P(4,N)*TANH(P(5,N)*ALOG(E/P(6,N)))
```

IF (E.LE.O) THEN

DO J=1,3

```
K=2*J+5
        DC=E/P(K,N)
        WC=EXP(-DC)
        C(J) = WC + (1.-WC)*DC**P(K+1,N)
       END DO
       PTOTAL = (A+B) *C(1) *C(2) *C(3)
       RETURN
       END
       SUBROUTINE LETAW(IZ, IA, ENERGY, CROSS_SECTION)
This subroutine computes the total inelastic cross sections
       of nuclides on protons. The formula is taken from:
       Letaw, J.R., Silberberg, R., and Tsao, C.H. 1983, Ap. J. Suppl.,
       51,271.
       REAL IA
       E=1.-.62*EXP(-ENERGY/200.)*SIN(10.9/ENERGY**.28)
       T=45.*IA**.7
       T=(1.+.016*SIN(5.3-2.63*LOG(IA)))*T*E
       IF(IZ.EQ.2) T=.8*T
       IF(IZ.EQ.4) T=(1.+.75*EXP(-ENERGY/75.))*T
       CROSS_SECTION=T
       RETURN
       END
```

```
SUBROUTINE CREME96 TRANSPORT(INPUT_FLUX,
                                    ELOWER, EUPPER, M, IZLO, IZUP,
                                    IPATH, UPATHO, TARGET, SHIELDFILE,
       &
                                    VERSION NUMBER, PROGRAM_CODE,
       æ
                                    OUTPUT FLUX)
  C
  This subroutine transports an input particle environment through a
  С
      specified thickness and type of shielding. It takes account both
  С
      ionization energy loss (dE/dx) as well as energy-dependent nuclear
  С
      fragmentation. The output is the particle environment (differential
  C
      fluxes vs. energy) inside the spacecraft, that is, 'behind' the specified
  С
      shielding. This routine includes many refinements over the old CREME
  С
      transport routine ("INSIDE"). Specifically:
  С
  C
      CREME96 TRANSPORT keeps track of projectile fragments; the old CREME
  C
      code ignored them. This routine also uses improved Silberberg, Tsao,
  C
      and Barghouty energy-dependent fragmentation cross-sections. Both of
  C
      these improvements can be important for thick shielding.
  С
  С
      At present CREME96_TRANSPORT only does aluminum shielding; future
  С
      versions will also offer transport through other shielding materials.
  С
  C
      CREME96_TRANSPORT is based on the "UPROP" code, as originally developed
  C
₫ c
      by John R. Letaw of Severn Communication Corp. under contract to
₽ C
      the Gamma & Cosmic Ray Astrophysics Branch of Naval Research Laboratory
C
      in 1989. Significant improvements and "bug-extermination" have been
Ī c
      provided by A.F. Barghouty of Roanoke College.
TI C
LI C
        IMPLICIT NONE
, in the second
        CHARACTER*12 TARGET
72
        CHARACTER*80 SHIELDFILE
ĿĿ
        INTEGER*4 MARR, NELM
TL.
        PARAMETER (MARR=5000, NELM=92)
REAL*4 INPUT FLUX (NELM, MARR), OUTPUT FLUX (NELM, MARR)
REAL*4 TEMP INPUT (NELM, MARR), TEMP_FLUX (NELM, MARR)
REAL*4 ELOWER, EUPPER, UPATHO, PATH, PSTEP, PSTEPMIN, PSTEPMAX
        REAL*4 PATHOLD, DELTA PATH, TEMP PATH
        INTEGER*4 M, N, NSP, IZLO, IZUP, IPATH, IULABEL
        REAL*4 UPATH, UUPATH, FRACSHLD
        INTEGER*4 VERSION NUMBER, PROGRAM CODE
        INTEGER*4 MAXSHIELD, NSHIELD, K, IELM, IARR
        PARAMETER (MAXSHIELD=500)
        DIMENSION UPATH (MAXSHIELD), FRACSHLD (MAXSHIELD)
        CHARACTER*5 UNITS_LABEL
        DIMENSION UNITS LABEL (4)
        DATA UNITS_LABEL/'g/cm2','mils ','cm ','!!!!!'/
        INTEGER*4 IENT
        DATA IENT/0/
   C
   C
        WRITE(6,9998)
    9998 FORMAT(1x,' TRANSPORT DRIVER calculation started.',
                  ' Please stand by.')
        CALL GET CREME96 VERSION (VERSION_NUMBER)
```

PROGRAM CODE=4

```
C
        Now set parametes for transport calculation.
  C
        Use recommended default:
  C
        Special version: turns off all nuclear fragmention:
  C
         IF (IENT.EQ.0) THEN
             IENT=1
             WRITE (6,9995)
             FORMAT(1x,'!!!! SPECIAL VERSION OF CREME96 TRANSPORT !!!!!',
   9995
                     /,' All nuclear fragmentation turned off!',
                     /,' NOTE: The user is responsible for tracking',
        &
                       ' to which output files this pertains!')
        ENDIF
        Use straight-ahead approximation; ignore energy spread of target fragments
  C
         (This takes a lot of time and generally has only very small effect.)
  С
         NSP=0
  C
         Set maximum & minimum PSTEP sizes allowed in transport
         PSTEPMIN=0.20
         PSTEPMAX=0.20
         IULABEL=IPATH+1
         IF (IULABEL.GT.4) IULABEL=4
IF (UPATHO.GT.O.O) THEN
             NSHIELD=1
             UPATH(1)=UPATHO
FRACSHLD(1)=1.00
             CALL UNLOAD SHIELDFILE (SHIELDFILE, NSHIELD, UPATH, FRACSHLD)
1....
         ENDIF
PATHOLD=0.0
N
IJ
DO 1000 K=1, NSHIELD
111
         WRITE(6,999) K, UPATH(K), UNITS_LABEL(IULABEL), FRACSHLD(K)
    999 FORMAT(1x,' SHIELDING BIN ', 14,' THICKNESS = ',F10.4,1x,A5,
                       FRACTION = ', F8.4)
         UUPATH=UPATH(K)
         Get shielding thickness (PATH) in g/cm2 and transport step size:
         CALL UNLOAD_PATH(IPATH, UUPATH, TARGET, PATH, PSTEPMIN, PSTEPMAX, PSTEP)
   C
         Now perform transport:
   \mathbf{C}
         IF (NSHIELD.EQ.1) THEN
         CALL UPROP96 (INPUT_FLUX,
                      ELOWER, EUPPER, M, IZLO, IZUP,
                      N, NSP, PATH, PSTEP, TARGET,
                       OUTPUT FLUX)
        ۶
   C
   C
         ELSE
   C
   C
         Modification 8-16-96 by AJT:
         To speed up calculations through thick shielding distributions,
   C
         allow output of one step to be input to the next step.
   C
```

¥

```
DELTA PATH=PATH-PATHOLD
         IF (DELTA PATH.LT.O.) DELTA_PATH=0.0
         DO 300 IELM=1, NELM
            DO 200 IARR=1, MARR
                   IF (K.EQ.1) THEN
                   TEMP INPUT (IELM, IARR) = INPUT FLUX (IELM, IARR)
                   TEMP PATH=PATH
                   ELSEIF (K.GT.1 .and. DELTA PATH.LT.PSTEP) THEN
                   TEMP INPUT (IELM, IARR) = INPUT_FLUX (IELM, IARR)
                   TEMP PATH=PATH
                   ELSEIF (K.GT.1 .and. DELTA_PATH.GE.PSTEP) THEN
                   TEMP INPUT (IELM, IARR) = TEMP FLUX (IELM, IARR)
                   TEMP PATH=DELTA PATH
     200
            CONTINUE
     300 CONTINUE
         CALL UPROP96 (TEMP INPUT,
                      ELOWER, EUPPER, M, IZLO, IZUP,
                      N, NSP, TEMP PATH, PSTEP, TARGET,
        &
                      TEMP FLUX)
         PATHOLD=PATH
        Now add to weighted sum:
        DO 500 IELM=1, NELM
            DO 400 IARR=1, MARR
               OUTPUT FLUX (IELM, IARR) = OUTPUT FLUX (IELM, IARR) +
                                       TEMP FLUX (IELM, IARR) *FRACSHLD(K)
        &
     400
            CONTINUE
     500 CONTINUE
         ENDIF
F.
    1000 CONTINUE
£
         WRITE(6,9999)
    9999 FORMAT(1x,' TRANSPORT_DRIVER calculation completed. Thank you.',
        & /,1x,' All fluxes are in units of particles/m2-s-sr-MeV/nuc',
              ' vs. energy in MeV/nuc.',
        &
        & /,1x,' Recommended next step: ',
        & /,5x,' LETSPEC',
               ' (RUN CREME96:LETSPEC DRIVER)'
              ' for heavy-ion induced SEUs;',
        & /,2x,' or PUP (RUN CREME96:PROTON_UPSET_DRIVER)',
              ' for proton-induced SEUs.')
         RETURN
         END
```

```
SUBROUTINE OUTPUT CREME96 DIFLET (LETMIN, LETMAX, L,
                                    IZLO, IZUP, EMINCUT, EMAXCUT, TARGET,
                                    VERSION NUMBER, PROGRAM CODE,
                                    INFILE,
                                    DIFSPEC,
                                    OUTFILE)
  C
        Routine for writing out the CREME96 differential LET Spectrum.
  С
  С
         IMPLICIT NONE
         REAL*4 LETMIN, LETMAX, EMINCUT, EMAXCUT, DIFSPEC
         DIMENSION DIFSPEC(1)
         INTEGER*4 L, IZLO, IZUP, LARR, K, NHEADER, NHEADERO, OUTUNIT
         DATA OUTUNIT/2/
         CHARACTER*12 TARGET
         CHARACTER*9 CREATION_DATE
         CHARACTER*8 CREATION TIME
         CHARACTER*80 INFILE, OUTFILE, DLTFILE
         INTEGER*4 VERSION NUMBER, PROGRAM CODE, STAT, CREME96_OPEN
   С
         FORMAT statements
         FORMAT (1X, 2 (1PE10.4, 2X), 3 (I5, 2X), A12, 16x, I4, 1x, I1)
   100
   150
         FORMAT(1x,A79)
         FORMAT((1X,6(1PE10.4,2X)))
200
210
         FORMAT((1X,6(E10.4,2X)))
         Open output file and write header
C C
THE C
  C
         First, modify name for DIFLET spectrum:
         DLTFILE='NULLFILE'
IJ
         DO K=2, LEN (OUTFILE)
1
               IF (OUTFILE(K:K) .EQ. '.') THEN
                   DLTFILE=OUTFILE(1:K-1)//'.DLT'
               ENDIF
         ENDDO
         IF (DLTFILE.EQ.'NULLFILE') DLTFILE=OUTFILE//'.DLT'
1
         OPEN (UNIT=OUTUNIT, STATUS='NEW', FILE='USER:'//DLTFILE)
         stat = creme96 open(dltfile, 'user', outunit, 'new')
         CALL DATE (CREATION DATE)
         CALL TIME (CREATION TIME)
         CALL CHECK HEADER LENGTH (INFILE, NHEADERO)
         NHEADER=NHEADER0+5
         WRITE (OUTUNIT, 990) NHEADER, DLTFILE (1:70),
                             VERSION NUMBER, PROGRAM CODE+1
    990 FORMAT(I3,1x,A70,I4,1x,I1)
         WRITE(OUTUNIT, 992) VERSION NUMBER, CREATION DATE, CREATION TIME
    992 FORMAT(1x,'%Created by CREME96:LETSPEC DRIVER Version', I4,
                    ' on ',A9,' at ',A8)
         WRITE (OUTUNIT, 993) IZLO, IZUP, LETMIN, LETMAX, L
    993 FORMAT(1x,'%ZMIN =',I3,' ZMAX =',I3,
               ' LETMIN = ',1PE8.2,' LETMAX = ',1PE8.2,
               ' MeV-cm2/g LBINS = ', I5)
         WRITE (OUTUNIT, 994) EMINCUT
    994 FORMAT(1x,'%EMINCUT = ',1PE8.2,' MeV/nuc')
         WRITE(OUTUNIT, 995) TARGET
    995 FORMAT(1x,'%TARGET MATERIAL = ',A12)
```

C Now copy header information from input file: WRITE(OUTUNIT,998) INFILE(1:45) 998 FORMAT(1x,'%Input File to LETSPEC_DRIVER: ',A45)

CALL COPY HEADERS (INFILE, NHEADERO, OUTUNIT)

C Finally, output differential LET spectrum:

WRITE (OUTUNIT, 100) LETMIN, LETMAX, L, IZLO, IZUP, TARGET,

VERSION_NUMBER, PROGRAM_CODE+1

WRITE (OUTUNIT, 100)

C Write flux to file.

WRITE(OUTUNIT,200) (DIFSPEC(K),K=1,L) CLOSE(OUTUNIT)

RETURN END

```
SUBROUTINE OUTPUT_CREME96_DOSE(INFILE,IZMIN,IZMAX,LETMIN,LETMAX,
                                   EMINCUT, EMAXCUT, TARGET, MODEL TYPE,
                                   VERSION NUMBER, PROGRAM CODE,
                                   DOSE PER SECOND, acc dose,
                                   OUTFILE)
  С
        Routine for writing out the CREME96 dose calculation.
  C
  С
        IMPLICIT NONE
        CHARACTER*80 INFILE, OUTFILE
        INTEGER*4 IZMIN, IZMAX, MODEL TYPE
        REAL*4 LETMIN, LETMAX, EMINCUT, EMAXCUT
        REAL*4 DOSE PER SECOND, acc dose
         INTEGER*4 NHEADER, NHEADERO, OUTUNIT
        DATA OUTUNIT/2/
         CHARACTER*12 TARGET
         CHARACTER*9 CREATION DATE
         CHARACTER*8 CREATION TIME
         INTEGER*4 VERSION NUMBER, PROGRAM CODE, STAT, CREME96_OPEN
        Open output file and write header
  С
         OPEN (UNIT=OUTUNIT, STATUS='NEW', FILE='USER:'//OUTFILE)
         stat = creme96 open(outfile, 'user', outunit, 'new')
         CALL DATE (CREATION DATE)
CALL TIME (CREATION_TIME)
CALL CHECK HEADER LENGTH (INFILE, NHEADERO)
         NHEADER=NHEADER0+6
         IF (MODEL TYPE.EQ.1 .or. MODEL TYPE.EQ.2) NHEADER=NHEADER+1
        WRITE (OUTUNIT, 990) NHEADER, OUTFILE (1:70),
                            VERSION NUMBER, PROGRAM CODE
990 FORMAT(I3,1x,A70,I4,I2)
         WRITE(OUTUNIT, 992) VERSION_NUMBER, CREATION DATE, CREATION TIME
992 FORMAT(1x,'%Created by CREME96:DOSE DRIVER Version', I4,
                   ' on ',A9,' at ',A8)
         IF(MODEL TYPE.EQ.0) WRITE(OUTUNIT, 900) DOSE PER SECOND, acc dose
IF(MODEL TYPE.EQ.1) WRITE(OUTUNIT, 901) DOSE PER SECOND, acc dose
         IF (MODEL TYPE.EQ.2) WRITE (OUTUNIT, 902) DOSE PER SECOND, acc dose
         IF(MODEL_TYPE.EQ.3) WRITE(OUTUNIT,903) DOSE_PER_SECOND,acc dose
    900 FORMAT(' %Average Dose = ',1PE13.6,' rad/sec = ',1PE13.6,
               ' krad/year')
    901 FORMAT(' %Worst-day average dose rate = ',1PE13.6,' rad/sec',
        & /,' %Event-Accumulated Dose = ',1PE13.6,' krad in 18.0 hours.')
    902 FORMAT(' %Worst-week average dose rate = ',E13.6,' rad/sec',
        & /,' %Event-Accumulated Dose = ',1PE13.6,' krad in 180.0 hours.')
    903 FORMAT(' %Peak SEP dose rate = '1P,E13.6,' rad/sec = ',
                  1PE13.6, 'krad/sec')
        &
         WRITE (OUTUNIT, 993) IZMIN, IZMAX, LETMIN, LETMAX
    993 FORMAT(1x,'%ZMIN =', I3,' ZMAX =', I3,
              ' LETMIN = ',1PE8.2,' LETMAX = ',1PE8.2,' MeV-cm2/g')
         WRITE (OUTUNIT, 994) EMINCUT, EMAXCUT
    994 FORMAT(1x,'%EMINCUT = ',1PE8.2,' EMAXCUT = ',1PE8.2,' MeV/nuc')
         WRITE (OUTUNIT, 995) TARGET
    995 FORMAT(1x,'%TARGET MATERIAL = ',A12)
         Now copy header information from input file:
         WRITE(OUTUNIT, 998) INFILE(1:45)
```

998 FORMAT(1x,'%Input File to DOSE_DRIVER: ',A45)
CALL COPY_HEADERS(INFILE,NHEADER0,OUTUNIT)

CLOSE (OUTUNIT)

RETURN END

```
SUBROUTINE OUTPUT_CREME96 FLUX(IZLO, IZHI, ELOWER, EUPPER,
                                      YEAR, IMODE, ITRANS,
                                      GTRANSFILE, TRAPDFILE,
                                      VERSION NUMBER, PROGRAM CODE,
                                      M, FLX, OUTFILE)
C
      Routine for writing the CREME96 particle environment file.
C
      Modified 7/29/96 to add tracking information to header.
С
     Modified 8/19/96 to add more detailed header information,
C
      per recommendation from Ed Petersen.
C
      IMPLICIT NONE
      INTEGER*4 IZLO, IZHI, J, K, M, OUTUNIT, STAT, CREME96 OPEN
      DATA OUTUNIT/2/
      INTEGER*4 NHEADER, NGTFLINES, NTRPLINES
      REAL*4 ELOWER, EUPPER
      CHARACTER*80 OUTFILE, ILINE
      CHARACTER*9 CREATION DATE
      CHARACTER*8 CREATION_TIME
      INTEGER*4 MARR, NELM
      PARAMETER (MARR=5000, NELM=92)
      REAL*4 FLX
      DIMENSION FLX (NELM, MARR)
      REAL*4 YEAR
      INTEGER*4 IMODE, ITRANS, VERSION_NUMBER, PROGRAM_CODE
      CHARACTER*80 GTRANSFILE, TRAPDFILE
      CHARACTER*12 TARGET
      DATA TARGET/'UNSHIELDED '/
     FORMAT statements
C
     FORMAT(1X,2(1PE10.4,2X),3(15,2X),A12,
100
              2X, OPF8.3, 1x, I2, 1x, I1, 1x, I4, 1x, I1)
     FORMAT (1x, A39, 1x, A39)
150
200
     FORMAT((1X,6(1PE10.4,2X)))
      Open output file and write header
C
       OPEN(UNIT=OUTUNIT,STATUS='NEW',FILE='USER:'//OUTFILE)
C
      stat = creme96 open(outfile,'user',outunit,'new')
      CALL DATE (CREATION DATE)
      CALL TIME (CREATION_TIME)
C
      Now prepare header for output file:
C
      NHEADER=3
       IF (ITRANS.EQ.0) NHEADER=NHEADER+1
       IF (ITRANS.GE.1) THEN
                   CALL CHECK_HEADER_LENGTH(GTRANSFILE,NGTFLINES)
                   NHEADER=NHEADER+2+NGTFLINES
       ENDIF
       IF (ITRANS.EQ.2) THEN
                   CALL CHECK HEADER LENGTH (TRAPDFILE, NTRPLINES)
                   NHEADER=NHEADER+1+NTRPLINES
       ENDIF
       WRITE(OUTUNIT, 990) NHEADER, OUTFILE(1:70),
             VERSION NUMBER, PROGRAM_CODE
  990 FORMAT(I3,1x,A70,I4,1x,I1)
       WRITE(OUTUNIT, 992) VERSION_NUMBER, CREATION_DATE, CREATION_TIME
```

```
' on ',A9,' at ',A8)
          Revised 9/14/96: Energy limits hardwired
          WRITE (OUTUNIT, 993) IZLO, IZHI, ELOWER, EUPPER, M
   C 993 FORMAT(1x,'%ZMIN =',I3,' ZMAX =',I3,
              ' EMIN = ',1PE10.4,' EMAX = ',1PE10.4,
              ' MeV/nuc MBINS =', I5)
         WRITE (OUTUNIT, 993) IZLO, IZHI
    993 FORMAT(1x,'%ZMIN =', I3,' ZMAX =', I3)
         IF (IMODE.EQ.0) THEN
            WRITE(OUTUNIT, 995) IMODE, YEAR
            FORMAT(1x,'%IMODE = ', I3,' SOLAR-QUIET MODE: YEAR = ', F10.4)
    995
         ELSEIF (IMODE.EQ.1) THEN
            WRITE (OUTUNIT, 996) IMODE
            FORMAT(1x,'%IMODE = ',I3,
                       ' WORST-DAY SOLAR ENERGETIC PARTICLE MODEL')
         ELSEIF (IMODE.EQ.2) THEN
            WRITE (OUTUNIT, 997) IMODE
            FORMAT(1x, '%IMODE = ', I3,
    997
                      ' WORST-WEEK SOLAR ENERGETIC PARTICLE MODEL')
        &
         ELSE
            WRITE(OUTUNIT,998) IMODE
FORMAT(1x,'%IMODE =', I3,' PEAK 5-MINUTE-AVERAGED FLUX')
         ENDIF
         IF (ITRANS.EQ.0) THEN
            WRITE (OUTUNIT, 999) ITRANS
            FORMAT(1x,'%ITRANS = ',I3,
    999
                      ' GEOSYNCH/NEAR-EARTH INTERPLANETARY FLUXES')
Ξ
         ELSEIF (ITRANS.EQ.1) THEN
            WRITE (OUTUNIT, 899) ITRANS, GTRANSFILE (1:40)
            FORMAT(1x,'%ITRANS = ',I3,
    899
' INSIDE MAGNETOSPHERE/NO TRAPPED FLUXES',
l.
                   /,1x,'%INPUT GEOMAGNETIC TRANSMISSION FILE:',1x,A40)
            CALL COPY_HEADERS (GTRANSFILE, NGTFLINES, OUTUNIT)
         ELSEIF (ITRANS.EQ.2) THEN
            WRITE (OUTUNIT, 898) ITRANS, GTRANSFILE (1:40)
            FORMAT(1x,'%ITRANS = ',I3,
    898
                       ' INSIDE MAGNETOSPHERE/TRAPPED PROTONS INCLUDED',
        &
                  /,1x,'%INPUT GEOMAGNETIC TRANSMISSION FILE:',1x,A40)
        &
            CALL COPY_HEADERS (GTRANSFILE, NGTFLINES, OUTUNIT)
             WRITE (OUTUNIT, 897) TRAPDFILE (1:40)
             FORMAT(1x,'%INPUT TRAPPED PROTON FILE: ',11x,A40)
    897
             CALL COPY_HEADERS (TRAPDFILE, NTRPLINES, OUTUNIT)
         ENDIF
         WRITE (OUTUNIT, 100) ELOWER, EUPPER, M, IZLO, IZHI,
                       TARGET, YEAR, IMODE, ITRANS,
                       VERSION NUMBER, PROGRAM_CODE
         WRITE (OUTUNIT, 100)
```

992 FORMAT(1x,'%Created by CREME96:FLUX DRIVER Version', I4,

DO J=IZLO,IZHI

WRITE(OUTUNIT,200) (FLX(J,K),K=1,M)

Skip line between elements. AJT 5/6/96

WRITE(OUTUNIT,100)

END DO

CLOSE (OUTUNIT)

RETURN END

С

```
IZLO, IZUP, EMINCUT, EMAXCUT, TARGET,
                                   VERSION NUMBER, PROGRAM CODE,
                                   INFILE,
                                   SPECT,
                                   OUTFILE)
  C
        Routine for writing out the CREME96 integral LET Spectrum.
  C
        Modified 7/29/96 to add header information.
  С
        Modified 8/19/96 to add more detailed header information, per
  С
        recommendation by Ed Petersen.
  C
  C
        IMPLICIT NONE
        REAL*4 LETMIN, LETMAX, EMINCUT, EMAXCUT
        INTEGER*4 L, IZLO, IZUP, LARR, K, NHEADER, NHEADERO, OUTUNIT
        DATA OUTUNIT/2/
        CHARACTER*12 TARGET
        CHARACTER*9 CREATION DATE
        CHARACTER*8 CREATION_TIME
        PARAMETER (LARR=1002)
        REAL*4 SPECT (LARR)
        CHARACTER*80 INFILE, OUTFILE
         INTEGER*4 VERSION_NUMBER, PROGRAM_CODE, STAT, CREME96_OPEN
        FORMAT statements
FORMAT(1X,2(1PE10.4,2X),3(I5,2X),A12,16x,I4,1x,I1)
  100
150
        FORMAT(1x, A79)
200
        FORMAT((1X,6(1PE10.4,2X)))
        Open output file and write header
LL C
į
        OPEN(UNIT=OUTUNIT,STATUS='NEW',FILE='USER:'//OUTFILE)
≝ . C
         stat = creme96 open(outfile,'user',outunit,'new')
1
         CALL DATE (CREATION DATE)
         CALL TIME (CREATION TIME)
Ų.
         CALL CHECK_HEADER_LENGTH(INFILE, NHEADER0)
         NHEADER=NHEADER0+5
         WRITE(OUTUNIT, 990) NHEADER, OUTFILE(1:70),
                            VERSION_NUMBER, PROGRAM_CODE
    990 FORMAT (I3, 1x, A70, I4, 1x, I1)
         WRITE(OUTUNIT, 992) VERSION NUMBER, CREATION DATE, CREATION TIME
    992 FORMAT(1x,'%Created by CREME96:LETSPEC_DRIVER Version',I4,
                   ' on ',A9,' at ',A8)
         WRITE (OUTUNIT, 993) IZLO, IZUP, LETMIN, LETMAX, L
    993 FORMAT(1x,'%ZMIN =',I3,' ZMAX =',I3,
              ' LETMIN = ',1PE8.2,' LETMAX = ',1PE8.2,
              ' MeV-cm2/g LBINS = ', I5)
         WRITE (OUTUNIT, 994) EMINCUT
    994 FORMAT(1x,'%EMINCUT = ',1PE8.2,' MeV/nuc')
         WRITE (OUTUNIT, 995) TARGET
    995 FORMAT(1x,'%TARGET MATERIAL = ',A12)
         Now copy header information from input file:
         WRITE (OUTUNIT, 998) INFILE (1:45)
    998 FORMAT(1x,'%Input File to LETSPEC_DRIVER: ',A45)
         CALL COPY_HEADERS (INFILE, NHEADERO, OUTUNIT)
         Finally, output integral LET spectrum:
```

SUBROUTINE OUTPUT CREME96 LETSPEC(LETMIN, LETMAX, L,

```
WRITE(OUTUNIT,100) LETMIN,LETMAX,L,IZLO,IZUP,TARGET,

VERSION_NUMBER,PROGRAM_CODE

WRITE(OUTUNIT,100)

Write flux to file.
```

CLOSE (OUTUNIT)

WRITE (OUTUNIT, 200) (SPECT (K), K=1, L)

RETURN END

С

```
COMMENT, IUNITS, MATERIAL,
                                    NBINS, XTHICK, XPROB,
                                    XMEAN, XRMS, TOTAL, ERRFLAG,
                                    VERSION NUMBER, PROGRAM CODE)
C
      IMPLICIT NONE
      CHARACTER*80 SHIELDFILE, COMMENT
      INTEGER*4 IUNITS, STAT, CREME96_OPEN
      CHARACTER*12 MATERIAL
      INTEGER*4 NBINS, VERSION NUMBER, PROGRAM_CODE
      REAL*4 XTHICK, XPROB, XMEAN, XRMS, TOTAL
      INTEGER*4 ERRFLAG
      DIMENSION XTHICK(1), XPROB(1)
      INTEGER*4 NHEADER
      INTEGER*4 OUTUNIT, IULABEL, K
      DATA OUTUNIT/2/
      CHARACTER*9 CREATION DATE
      CHARACTER*8 CREATION_TIME
      CHARACTER*5 UNITS LABEL
      DIMENSION UNITS LABEL(4)
      DATA UNITS LABEL/'g/cm2', 'mils ', 'cm ', '!!!!!'/
      Open output file and write header
      OPEN(UNIT=OUTUNIT,STATUS='NEW',FILE='USER:'//SHIELDFILE)
      stat = creme96_open(shieldfile,'user',outunit,'new')
      CALL DATE (CREATION DATE)
      CALL TIME (CREATION TIME)
      NHEADER=4
      WRITE (OUTUNIT, 990) NHEADER, SHIELDFILE (1:70),
                          VERSION NUMBER, PROGRAM CODE
 990 FORMAT(I3,1x,A70,I4,1x,I1)
      WRITE (OUTUNIT, 991) VERSION NUMBER, CREATION DATE, CREATION TIME
 991 FORMAT(1x,'%Created by CREME96:SHIELDFILE_DRIVER Version', I4,
                 ' on ',A9,' at ',A8)
      WRITE (OUTUNIT, 992) COMMENT (1:78)
 992 FORMAT(1X,'%',A78)
      IULABEL=IUNITS+1
      IF (IULABEL.GT.4) IULABEL=4
      WRITE(OUTUNIT, 993) UNITS LABEL(IULABEL), MATERIAL, NBINS
 993 FORMAT(1x,'%Shielding thicknesses: in ',A5,4x,A12,
                                           3x, 'NBINS = ', I5)
      IF (ERRFLAG.EQ.0) WRITE(OUTUNIT,994) XMEAN, XRMS
  994 FORMAT(1x'%MEAN = ',E13.6,' RMS = ',E13.6)
      IF (ERRFLAG.EQ.1) WRITE (OUTUNIT, 995) XMEAN, XRMS, TOTAL
  995 FORMAT(1x'%MEAN = ',E13.6,' RMS = ',E13.6,
                 ' Sum Before Renorm = ',E13.6)
      WRITE (OUTUNIT, 996) IUNITS
  996 FORMAT(I2)
       DO 1000 K=1, NBINS
          WRITE (OUTUNIT, 999) XTHICK (K), XPROB (K)
          FORMAT (1x, E13.6, 5x, E13.6)
  999
 1000 CONTINUE
       CLOSE (OUTUNIT)
```

RETURN

SUBROUTINE OUTPUT SHIELDFILE (SHIELDFILE,

```
SUBROUTINE OUTPUT TRANSPORTED FLUX (IZLO, IZHI, ELOWER, EUPPER,
                                       IPATH, UPATH, TARGET,
                                       SHIELDFILE, INFILE,
                                       VERSION_NUMBER, PROGRAM CODE,
                                       M, FLX, OUTFILE)
C
      Routine for writing the CREME96 transported particle environment file.
C
      Modified 7/29/96 to add tracking information to header.
C
      Modified 8/19/96 to add more detailed header information, per
С
      recommendation by Ed Petersen.
C
C
      IMPLICIT NONE
      INTEGER*4 IZLO, IZHI, J, K, M, NHEADER, NHEADER0, IULABEL
      INTEGER*4 OUTUNIT, STAT, CREME96_OPEN
      DATA OUTUNIT/2/
      REAL*4 ELOWER, EUPPER
      CHARACTER*80 OUTFILE, ILINE
      CHARACTER*9 CREATION DATE
      CHARACTER*8 CREATION TIME
      CHARACTER*5 UNITS_LABEL
      DIMENSION UNITS LABEL (4)
      DATA UNITS_LABEL/'g/cm2','mils ','cm ','!!!!!'/
      INTEGER*4 MARR, NELM
      PARAMETER (MARR=5000, NELM=92)
      REAL*4 FLX
      DIMENSION FLX (NELM, MARR)
    REAL*4 UPATH
    INTEGER*4 IPATH, IDUM, VERSION_NUMBER, PROGRAM_CODE, NSHDLINES
      DATA NSHDLINES/0/
     CHARACTER*80 INFILE, SHIELDFILE
      CHARACTER*12 TARGET
      DATA IDUM/0/
     FORMAT statements
     FORMAT(1X,2(1PE10.4,2X),3(I5,2X),A12,
100
              2X, OPF8.3, 1x, I2, 1x, I1, 1x, I4, 1x, I1)
     FORMAT (1x, A39, 1x, A39)
150
200
     FORMAT((1X,6(1PE10.4,2X)))
      Open output file and write header
C
       OPEN (UNIT=OUTUNIT, STATUS='NEW', FILE='USER:'//OUTFILE)
C
       stat = creme96_open(outfile,'user',outunit,'new')
       CALL DATE (CREATION DATE)
       CALL TIME (CREATION_TIME)
C
      CALL CHECK_HEADER_LENGTH(INFILE,NHEADER0)
      NHEADER=NHEADER0+4
                                              ') THEN
       IF (SHIELDFILE(1:12).NE.'
           CALL CHECK HEADER LENGTH (SHIELDFILE, NSHDLINES)
           NHEADER=NHEADER+NSHDLINES
       ENDIF
       WRITE (OUTUNIT, 990) NHEADER, OUTFILE (1:70),
                           VERSION NUMBER, PROGRAM CODE
  990 FORMAT(I3,1x,A70,I4,1x,I1)
```

×

man the min

```
WRITE(OUTUNIT, 992) VERSION_NUMBER, CREATION_DATE, CREATION_TIME
992 FORMAT(1x,'%Created by CREME96:TRANSPORT DRIVER Version', I4,
               ' on ',A9,' at ',A8)
     WRITE (OUTUNIT, 993) IZLO, IZHI, ELOWER, EUPPER, M
993 FORMAT(1x,'%ZMIN =', I3,' ZMAX =', I3,
          ' EMIN = ',1PE10.4,' EMAX = ',1PE10.4,
          ' MeV/nuc MBINS =', I5)
     IULABEL=IPATH+1
     IF (IULABEL.GT.4) IULABEL=4
                                            ') THEN
     IF (SHIELDFILE(1:12).NE.'
        WRITE (OUTUNIT, 997) SHIELDFILE (1:50)
        FORMAT(1x,'%Shielding distribution: ',A50)
997
        CALL COPY_HEADERS (SHIELDFILE, NSHDLINES, OUTUNIT)
     ELSE
        WRITE(OUTUNIT, 996) UPATH, UNITS_LABEL(IULABEL), TARGET
        FORMAT (1x, '%Thickness = ', F10.4, 1x, A5, 5x, A12)
996
     ENDIF
     Now copy header information from input file:
     WRITE (OUTUNIT, 998) INFILE (1:45)
998 FORMAT(1x,'%Input File to TRANSPORT_DRIVER: ',A45)
     CALL COPY_HEADERS (INFILE, NHEADERO, OUTUNIT)
     Finally, output transported spectra:
     WRITE(OUTUNIT, 100) ELOWER, EUPPER, M, IZLO, IZHI,
                   TARGET, UPATH, IPATH, IDUM,
                   VERSION NUMBER, PROGRAM_CODE
     WRITE (OUTUNIT, 100)
     Write fluxes to file.
     DO J=IZLO, IZHI
       WRITE (OUTUNIT, 200) (FLX (J, K), K=1, M)
       Skip line between elements. AJT 5/6/96
       WRITE (OUTUNIT, 100)
     END DO
     CLOSE (OUTUNIT)
     RETURN
     END
```

```
SUBROUTINE PARTIALS (ENERGY, IZ, IA, NAME, MEAN FREE_PATH,
     & NSP, ELOSS, NORMALIZATION)
  _____
       This program computes the charge changing mfps
        for the nuclide (IZ,IA) on material NAME at ENERGY MeV/N.
        The target nuclides are (JZ, JA) and the products are (KZ, KA)
ļ-----
        INCLUDE 'CREME96: ZCOMMON.CMN'
        CHARACTER*12 NAME
        REAL NA(28), IADJ(28), NASPM(28), DENS, ETAD
        INTEGER NZ(28), IGAS, NAS
        COMMON /TBLOCK/DENS, ETAD, IGAS, NAS,
                      NZ, NA, IADJ, NASPM,
     &
                      NTOTAL, AVGZ, AVGZ2, AVGA, AVGI
     &
        REAL MEAN FREE PATH(100), CR(100), ELOSS(100), NORMALIZATION
        REAL IA, JA, KA
        DATA AVOGADRO/6.022045E23/ ! particles/mole
        CALL ZTARGET (NAME)
        FACT=1.E27*AVGA/AVOGADRO
        NORMALIZATION=0.
        DO KZ=3, IZ+1
          MEAN FREE PATH(KZ)=0.
          DO K=KZ,3*KZ
            KA=FLOAT(K)
            DO L=1, NAS
              JZ=NZ(L)
              JA=NA(L)
C..... The following modulates the projectile energy by the number of
C......TARGET participants, July 1992:
              IF (NINT (JA), EQ.1) THEN
                E=AMIN1 (ENERGY*JA, 10000.)
              ELSE
                 AP=IA
                 AT=JA
                 CALL GLAUBER (AP, AT, AP_P, AT_P)
                 PART=AT P+AP P
                 DELTA=IA-KA
                 A EFF=AT P*(1.+TANH((DELTA-PART)/PART))
                 E=AMIN1(ENERGY*A EFF, 10000.)
              END IF
C
              CALL YIELDX (IZ, NINT (IA), KZ, NINT (KA), AMAX1 (E, 100.), S)
              IF (KZ.EQ.4.AND.K.EQ.8) S=0.
              IF (KZ.EQ.5.AND.K.EQ.9) S=0.
              IF (S.LT.1.E-4) S=0.
              CALL SCALER(IZ, NINT(IA), JZ, NINT(JA), KZ, NINT(KA),
              AMAX1 (ENERGY, 100.), SC)
              MEAN_FREE_PATH(KZ)=MEAN_FREE_PATH(KZ)+S*SC*NASPM(L)
              NORMALIZATION=NORMALIZATION+S*SC*NASPM(L)*KZ
            END DO
          END DO
          MEAN FREE PATH(KZ)=MEAN FREE_PATH(KZ)/FACT/NTOTAL
        END DO
        MEAN FREE PATH(2)=0.
        DO K2=1,3
          IF (K2.EQ.1) KZ=2
          IF (K2.EQ.2) KZ=4
          IF (K2.EQ.3) KZ=5
```

```
IF (K2.EQ.1) K=4
             IF (K2.EQ.2) K=8
             IF (K2.EQ.3) K=9
             KA=FLOAT(K)
             DO L=1, NAS
               JZ=NZ(L)
               JA=NA(L)
   C.....The following modulates the projectile energy by the number of
   C......TARGET participants, July 1992:
                  IF(NINT(JA).EQ.1) THEN
                    E=AMIN1 (ENERGY*JA, 10000.)
                  ELSE
                    AP=IA
                    AT=JA
                    CALL GLAUBER (AP, AT, AP_P, AT_P)
                    PART=AT P+AP P
                    DELTA=IA-KA
                    A EFF=AT P*(1.+TANH((DELTA-PART)/PART))
                    E=AMIN1 (ENERGY*A_EFF, 10000.)
   С
                CALL YIELDX (IZ, NINT (IA), KZ, NINT (KA), AMAX1 (E, 100.), S)
                IF (KZ.EQ.4.OR.KZ.EQ.5) S=S*2.
                IF (S.LT.1.E-4) S=0.
                CALL SCALER (IZ, NINT (IA), JZ, NINT (JA), KZ, NINT (KA),
                AMAX1 (ENERGY, 100.), SC)
                MEAN_FREE_PATH(2)=MEAN_FREE_PATH(2)+S*SC*NASPM(L)
                NORMALIZATION=NORMALIZATION+S*SC*NASPM(L)*2.
             END DO
           END DO
           MEAN FREE PATH(2)=MEAN_FREE_PATH(2)/FACT/NTOTAL
           NORMALIZATION=NORMALIZATION/FACT/NTOTAL
<u>End</u>
×
   C.....Energy-loss calculations: Sept. 1993
           IF(NSP.EQ. 0) RETURN
DO KZ=1, IZ+1
ELOSS (KZ) = 0.
              CR(KZ) = 0.
             DO K=KZ,3*KZ
                KA=FLOAT(K)
                DO L=1, NAS
                  JZ=NZ(L)
                  JA=NA(L)
   C.....The following modulates the projectile energy by the number of
   C......TARGET participants, July 1992:
                  IF (NINT (JA) . EQ.1) THEN
                     E=AMIN1 (ENERGY*JA, 10000.)
                  ELSE
                     AP=IA
                     AT=JA
                     CALL GLAUBER (AP, AT, AP P, AT P)
                     PART=AT P+AP P
                     DELTA=IA-KA
                     A_EFF=AT_P*(1.+TANH((DELTA-PART)/PART))
                     E=AMIN1 (ENERGY*A EFF, 10000.)
                  END IF
   C
```

CALL YIELDX (IZ, NINT (IA), KZ, NINT (KA), AMAX1 (E, 100.), S)

```
IF (KZ.EQ.4.AND.K.EQ.8) S=0.
                  IF (KZ.EQ.5.AND.K.EQ.9) S=0.
                  IF (S.LT.1.E-4) S=0.
                  CALL SCALER (IZ, NINT (IA), JZ, NINT (JA), KZ, NINT (KA),
                 AMAX1 (ENERGY, 100.), SC)
        &
                  CALL E LOSS (IZ, NINT (IA), JZ, NINT (JA), KZ, NINT (KA),
                              ENERGY, dKE, SigmaKE)
                  dKE=dKE+0.*SigmaKE
                  CR(KZ) = CR(KZ) + S*SC*NASPM(L) / NTOTAL
                  IF (KZ.EQ.1) THEN
                     ELOSS (KZ) = ELOSS (KZ) + dKE* (REAL (KA) /6.) *NASPM (L)
                                 /NTOTAL
        &
                  ELSE
                     ELOSS(KZ)=ELOSS(KZ)+dKE*S*SC*NASPM(L)/NTOTAL
                  END IF
               END DO
             END DO
                  IF (CR (KZ) .NE.O.AND.KZ.GT.1) THEN
                    ELOSS (KZ) = ELOSS (KZ) / CR (KZ)
                  END IF
           END DO
   C
           RETURN
           END
J.
           SUBROUTINE SCALER (IZ1, IA1, IZ2, IA2, JZ, JA, E, SC)
           DATA IENT/0/
           IF (IENT.EQ.0) THEN
TU
             IENT=1
  C
              TYPE *,' '
              TYPE *,'
                          ^^^ Using STB scaling algorithm-1992 ^^^'
С
              TYPE *,' '
           END IF
3
           SC = 1.
1
           IF (IA2.EQ.1) RETURN
SL = 1.
S1 = 1.
Ŀ£
           SD = 1.
           Z1 = IZ1
A1 = IA1
           Z2 = IZ2
           A2 = IA2
           Z = JZ
           A = JA
           E1 = E/1000.
   С
           SC = 1.6 + 0.07*A2**(2./3.)
   C......New scaling algorithm: July 1992
           SC=ASYMM (A1, A2)
   С
            FA = (1.0 + (A1/120.) *AMIN1(E1,2.)/2.)/(1.+A1/120.)
            IF (JZ.LE.5) SL = (1. + .4*(1.+.02*(Z1/Z)**2)*(1.-1.5*Z/Z1))*FA
            IF (A .LT.A1/2 ...AND. JZ.GT.5) SD = 3.*EXP(-(2.*A/A1))*FA
            IF (IA1-JA.EQ.\ 1) S1 = (1. + .0014*Z1*Z2**(1.8-.005*Z2))/SC
            SC = SC*SL*S1*SD
            END
            SUBROUTINE GLAUBER (AP, AT, AP P, AT P)
   C
            calculates (average) number of proj. and target participants;
```

```
according to Glauber theory.
C
С
        DATA PI,R0/3.14159,1.36/
        DATA P13, P23/0.33333, 0.66667/
С
        AP P=AP * AT**P23 / (AP**P13+AT**P13)**2
        AT_P=AT * AP**P23 / (AP**P13+AT**P13)**2
        RETURN
        END
C
        FUNCTION ASYMM (AP, AT)
        calculates asymmetry and participant contributions:
С
        DATA PI,R0/3.14159,1.36/
        DATA P13, P23/0.33333, 0.66667/
С
        CALL GLAUBER (AP, AT, AP_P, AT_P)
C
        EXPO=EXP(-(AP-AT)/(AT+AP))
        RHO=(AP-1.)/(AP+1.)
                                          !Normalization constant.
        CONST=EXP(-RHO)
        ASYMM=CONST*(AP/AP_P)*EXPO
        RETURN
        END
```

C

C С

> C С C

> С С

> C

С

C С

С С

С

C

С

C

С

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C

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C

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С

С С C

Module: PC ROUTINES

C Logical Names and Environment Variables serve the same purpose, C but are handled differently, on the two CREME96 platforms (VAX and C PC respectively). There are also differences between the two file C OPEN statements. To enable platform independance where fully C specified filenames and where file opens are used in the higher С level CREME96 code, two versions exist of the routines to perform С these tasks. When an executable is being created, it is the С responsibility of the person performing the link to ensure that the С

appropriate set of routines is used for the current build.

Two platform-DEPENDANT routines exist:

creates fully-specified filename CREME96 FULL FILENAME performs a file OPEN on full filename CREME96 OPEN gives advice when file not found. SHOW DIRECTORY

These routines reside in the following 2 physical files:

used for a VAX build VAX ROUTINES.FOR used for a PC build (this file) PC ROUTINES.FOR

integer function creme96_open(filename,path,unit,status)

The non-fully specified name of the target file. FILENAME:

Contains the DOS Environment Variable pointing to PATH:

the directory where file does, or will exist.

The logical unit to be associated with the file. UNIT:

Must be defined at the time of the function call

(one will not be assinged by this routine).

Contains either OLD, for existing file, or STATUS:

NEW, to create a file.

Calling example:

STAT = creme96 open('input.dat','creme96',inunit,'old')

Success is indicated by a ZERO return value. Otherwise, the return value will contain the FORTRAN error code.

IMPLICIT NONE

filename, file, full filename, line character*80

character*10 path unit, ios integer

character*7

status

```
file = full_filename(filename,path)
       if (status(1:1) .eq. 'o' .or. status(1:1) .eq. 'O') then
          Old files are only opened for READ (no APPEND in CREME).
С
          Any file opened for READ will be opened SHARED.
C
          OPEN(UNIT=unit, file=file, status='old',
          mode='read',share='denywr',iostat=ios,err=199)
c DEBUG
          read(unit, 99)line
С
          format(a80)
c99
          write(*,*)'First line in file: ',line
С
       else
          New file to be created. WRITE and NOSHARE are default. On the PC,
С
          we must open with REPLACE instead of NEW, in case a file already
C
          exists of this name (as it is our intention to write over it). If
С
          one doesn't exist, REPLACE acts the same as NEW.
C
          OPEN (UNIT=unit, file=file, status='replace',
          iostat=ios,err=199)
c DEBUG
          write(*,*)'Writing test line to new file...'
С
          write(unit,*)'Test line'
С
        endif
        creme96 open file = ios
199
        return
        end
С
        *******************
C*
С
        character*80 function full_filename(filename,path)
        use msflib
        IMPLICIT NONE
        character*80
                       filename, dir
                       path
        character*10
                       lendir
        integer
        The variable PATH contains the name of the environment variable
С
        which in turn points to the directory path of the target file.
С
        The function GETENVQQ will translate this environment variable
С
        into the program variable DIR.
C
        lendir = getenvqq(path(1:len_trim(path)),dir)
        full filename = dir(1:lendir)//'\'//filename
        return
        end
        ********************
С
       SUBROUTINE SHOW DIRECTORY (JFILETYPE)
C
C
       PC version.
```

--

<u>=</u>

S.

I

```
C
         The VAX version of this routine uses LIB$SPAWN to echo back a copy
  C
         of the user's directory when this routine becomes activated (because
  C
         the requested file was not found, etc. ) However, no comparable
         capability exists on the PC. We therefore just print out statements
  С
         which recommend that the user open another window and check his/her
  C
  C
         directory.
  C
         INTEGER*4 JFILETYPE
         IF (JFILETYPE.EQ.O) THEN
          WRITE(6,9010)
         FORMAT(1x,' Please open another window and',
       & ' check the directory of your current USER area:')
         ELSEIF (JFILETYPE.EQ.1) THEN
          WRITE(6,9011)
         FORMAT(1x,' Please open another window and',
    9011
       & ' check the directory of your *.tr* files.')
         ELSEIF (JFILETYPE.EQ.2) THEN
          WRITE(6,9012)
         FORMAT(1x,' Please open another window and',
        & ' check the directory of your *.gt* files.')
         ELSEIF (JFILETYPE.EQ.3) THEN
          WRITE(6,9013)
         FORMAT(1x,' Please open another window and',
       & ' check the directory of your particle flux files:',/,
& ' *.flx, *.tfx, *.tr*')
         ELSEIF (JFILETYPE.EQ.4) THEN
WRITE(6,9014)
         FORMAT(1x,' Please open another windw and',
.
       & ' check the directory of your particle flux files:',/,
& ' *.tfx, *.flx, *.tr*')
Ų.
ELSEIF (JFILETYPE.EQ.5) THEN
ű
          WRITE(6,9015)
         FORMAT(1x,' Please open another window and',
        & ' check the directory of your *.LET files.')
         ELSEIF (JFILETYPE.EQ.6) THEN
          WRITE (6, 9016)
         FORMAT(1x,' Please open another winow and',
        & ' check the directory of your *.DLT files.')
         ELSEIF (JFILETYPE.EQ.7) THEN
          WRITE(6,9017)
         FORMAT(1x,' Please open another window and',
        & ' check the directory of your *.SHD files.')
         ELSEIF (JFILETYPE.EQ.8) THEN
          WRITE (6,9018)
         FORMAT(1x,' Please open another window and',
        & ' check the directory of your *.XSD files.')
          ENDIF
          WRITE (6, 9999)
```

C

```
SUBROUTINE PROTON_UPSETS(PROTON_FILE, IPARAM, PARAMS,
                                    XSECT FILE, NBITS, IENTER,
       &
                                    SEU_RATE, DAY_RATE, PERSECOND, PERDAY)
       &
  С
          Subroutine for performing proton SEU evaluation:
  C
          Inputs: PROTON FILE = file containing proton differential flux
  C
                                  (in protons/m2-s-sr-MeV) vs. energy (in MeV)
  C
                                = 1,2,4, indicating cross-section model
                    IPARAM
  С
                                  1 = Bendel 1-parameter
  С
                                  2 = Bendel 2-parameter
  C
                                  4 = Weibull
  C
                                  0 = table
  C
                                    = array containing cross-section parameters
                    PARAMS (4)
  С
                    XSECT FILE = file containing cross-section table.
  C
                               = steering flag, for multiple calculations with
                    IENTER
  C
                                 the same proton spectrum.
  C
  C
          Outputs: SEU_RATE
                               in SEUs/s/bit
  C
                     DAY RATE in SEUs/bit/day
  С
                     PERSECOND in SEUs/device/second
  C
                                 in SEUs/device/day
                     PERDAY
  С
  С
                          Allan J. Tylka
  С
          Written by:
⊒ c
                          Code 7654
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TU C
          Last update: 20 August 1996
C
LI C
ini C-
  С
           IMPLICIT NONE
INTEGER*4 NBINS, NPTS, IPARAM, IENTER
T.
           REAL*4 EN, FLUX, PARAMS, XSECT, NBITS,
                  SEU_RATE, DAY_RATE, PERSECOND, PERDAY
CHARACTER*80 PROTON_FILE, XSECT_FILE
ũ
           PARAMETER (NBINS=5000)
           DIMENSION EN(NBINS), FLUX(NBINS), XSECT(NBINS)
           DIMENSION PARAMS (4)
           WRITE(6,9998)
           FORMAT(1x,' PROTON_UPSET_DRIVER calculation started.',
    9998
                   ' Please stand by.')
           SEU RATE=0.0
   C
           On first entry, get proton spectrum:
   C
           IF (IENTER.EQ.1) THEN
           CALL UNLOAD PROTON SPECTRUM(PROTON_FILE, EN, FLUX, NPTS)
```

ENDIF

```
Evaluate proton SEU cross-section at these energy values:
С
        CALL EVALUATE_SEU_CROSS_SECTION(EN,NPTS,IPARAM,PARAMS,
                                        XSECT_FILE, XSECT)
     &
C
        Calculate SEU rate:
        CALL INTEGRATE_PROTON_UPSETS(NPTS,EN,FLUX,XSECT,SEU_RATE)
        IF (SEU_RATE.LT.0.) THEN
            WRITE(6,999) SEU_RATE
            FORMAT(1x,' ERROR in PROTON_UPSETS: SEU RATE = ',E13.5)
 999
            SEU RATE=0.0
        ENDIF
        CALL CALC_SEU_RATE(NBITS,SEU_RATE,DAY_RATE,PERSECOND,PERDAY)
        WRITE(6,9999)
        FORMAT(1x,' PROTON_UPSET_DRIVER calculation completed. ')
 9999
        RETURN
        END
```

```
PROGRAM PROTON UPSET DRIVER
         IMPLICIT NONE
         REAL*4 NBITS, PARAMS, SEU_RATE, DAY_RATE, PERSECOND, PERDAY
         REAL*4 XDUM
         INTEGER*4 IPARAM, IREPEAT, IENTER
         DIMENSION PARAMS (4)
         CHARACTER*80 PROTON_FILE, XSECT_FILE, REPORT_FILE
         CHARACTER*40 DEVICE_LABEL
         INTEGER*4 IERR
         DATA IERR/0/
         IENTER=1
         CONTINUE
   10
         CALL INITIALIZE PROTON UPSETS (PROTON FILE, NBITS,
                                         IPARAM, PARAMS, XSECT FILE, IENTER,
                                         DEVICE LABEL, REPORT_FILE)
        &
         CALL PROTON_UPSETS(PROTON_FILE, IPARAM, PARAMS, XSECT_FILE,
                              NBITS, IENTER,
        &
                              SEU_RATE, DAY_RATE, PERSECOND, PERDAY)
        &
          CALL PROTON UPSET REPORT (PROTON_FILE, NBITS,
                                    IPARAM, PARAMS, XSECT FILE, IENTER,
        &
                                    DEVICE LABEL, REPORT FILE,
        &
                                    SEU RATE, DAY_RATE, PERSECOND, PERDAY)
&
              CONTINUE
    9100
              CALL RETRY INPUT (IERR)
              WRITE(6,9200)
              FORMAT(//,' Repeat SEU rate calculation with different',
- Hand
    9200
                         ' device characteristics? (1=yes,0=no)')
READ(*,*,ERR=9100,IOSTAT=IERR) IREPEAT
              IF (IREPEAT.EQ.1) THEN
                  IENTER=IENTER+1
                   GOTO 10
              ENDIF
          WRITE(6,9600)
    9600 FORMAT(1x,' Proton Upset calculations finished.')
          STOP
          END
```

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SUBROUTINE PROTON_UPSET_REPORT(PROTON_FILE, NBITS,
                                IPARAM, PARAMS, XSECT FILE, IENTER,
   æ
                                DEVICE LABEL, REPORT FILE,
   &
                                SEU RATE, DAY RATE, PERSECOND, PERDAY)
   æ
      IMPLICIT NONE
      REAL*4 NBITS, PARAMS, SEU RATE, DAY RATE, PERSECOND, PERDAY
      INTEGER*4 IPARAM, IENTER, OUTUNIT, VERSION NUMBER, NHEADERO, K
      INTEGER*4 NHEADER, PROGRAM CODE, STAT, CREME96 OPEN
      DATA OUTUNIT/2/
      DIMENSION PARAMS (4)
      CHARACTER*80 PROTON FILE, XSECT FILE, REPORT FILE
      CHARACTER*40 DEVICE LABEL
      CHARACTER*9 CREATION DATE
      CHARACTER*8 CREATION TIME
      PROGRAM CODE=9
      IF (IENTER.EQ.1.and.REPORT_FILE.NE.'NULLFILE') THEN
        OPEN(UNIT=OUTUNIT, FILE='USER:'//REPORT_FILE, STATUS='NEW')
       stat = creme96_open(report_file, 'user', outunit, 'new')
       CALL DATE (CREATION DATE)
       CALL TIME (CREATION TIME)
       CALL GET CREME96 VERSION (VERSION_NUMBER)
       CALL CHECK HEADER LENGTH (PROTON FILE, NHEADERO)
       NHEADER=NHEADER0+2
       WRITE (OUTUNIT, 991) NHEADER, REPORT_FILE (1:70),
                           VERSION_NUMBER, PROGRAM_CODE
991
       FORMAT(I3,1x,A70,I4,I2)
       WRITE(OUTUNIT, 992) VERSION_NUMBER, CREATION_DATE, CREATION_TIME
       FORMAT(1x,'%Created by CREME96:PROTON UPSET DRIVER Version', I4,
992
               ' on ',A9,' at ',A8)
    &
       Now copy header information from input file:
       WRITE(OUTUNIT,993) PROTON_FILE(1:40)
       FORMAT(1x,'%Input Proton Spectrum File: ',A40)
       CALL COPY HEADERS (PROTON_FILE, NHEADERO, OUTUNIT)
      IF (REPORT FILE.NE.'NULLFILE') THEN
       WRITE (OUTUNIT, 994) IENTER, DEVICE_LABEL
       FORMAT (/,1x,' REPORT NO. ',14,': ',2x,A40)
994
       IF (IPARAM.EQ.0) WRITE(outunit, 980) IPARAM, XSECT_FILE(1:75)
       IF (IPARAM.EQ.1) WRITE(outunit, 981) IPARAM, PARAMS(1)
       IF (IPARAM.EQ.2) WRITE(outunit, 982) IPARAM, PARAMS(1), PARAMS(2)
       IF (IPARAM.EO.4) WRITE (outunit, 984) IPARAM, (PARAMS(K), K=1,4)
       WRITE(outunit,996) NBITS
       FORMAT(1x,') Number of bits = ',E13.5)
996
       FORMAT(1x,' CROSS-SECTION INPUT ', 13,' FROM FILE: ',
980
            /,5x,A75)
       FORMAT(1x,' CROSS-SECTION INPUT', 13,
981
                   BENDEL 1-PARAMETER = ',E13.5)
       FORMAT(1x,' CROSS-SECTION INPUT', I3,
982
                 ' BENDEL 2-PARAMETERS A,B = ',2E13.5)
       FORMAT(1x,' CROSS-SECTION INPUT', 13,
984
                 ' WEIBULL FIT: ',
            /,5x,' ONSET = ',F9.3,' MeV',
            /,5x,' WIDTH = ',F9.3,' MeV',
```

```
SUBROUTINE RANGE(E,N,ZO,A1,NAME,R)
       ******************
С
          THIS PROGRAM TABULATES THE RANGE OF NUCLIDE (Z0,A1) IN
С
          A STOPPING MEDIUM 'NAME' AT ENERGIES GIVEN IN THE ARRAY
С
C
          E IN MeV/nucleon.
       *******************
С
       CHARACTER*12 NAME
       DIMENSION E(N), R(N), GX(4), GA(4)
       COMMON AVGZ, AVGZ2, AVGA, AVGI
       DATA NGAUSS/8/
C
       DATA GX/1.96028986,1.79666648,1.52553241,1.18343464,
C
               0.81656536,0.47446759,0.20333352,0.03971014/
C
       DATA GA/0.10122854,0.22238103,0.31370665,0.36268378,
С
               0.36268378,0.31370665,0.22238103,0.10122854/
C
       DATA NGAUSS/4/
       DATA GX/1.86113631,1.33998104,0.66001896,0.13886369/
       DATA GA/0.34785485,0.65214515,0.65214515,0.34785485/
       SS=STPOW(E(1),Z0,A1,NAME)
       JTEST=0
       DO J=1,N
         IF (E(J).LE.O.) THEN
           R(J)=0.
         ELSE
            IF (JTEST.EQ.0) THEN
             ELAST=0.
             RLAST=0.
              JTEST=1
            ELSE
              ELAST=E(J-1)
              RLAST=R(J-1)
            ENDIF
            DE = (E(J) - ELAST)/2.
            R(J)=0.
            DO K=1, NGAUSS
              STEMP=STPOW(ELAST+DE*GX(K), Z0, A1, NAME)
              IF (STEMP.GT.O.) R(J)=R(J)+GA(K)/STEMP
            END DO
            R(J) = DE * R(J) + RLAST
          ENDIF
        END DO
        RETURN
        END
```

```
SUBROUTINE RETRY INPUT (IERR)
  С
         NOTE: A non-zero input value of IERR will be re-set to zero
  С
         by this routine.
  С
  C
         IMPLICIT NONE
         INTEGER*4 IERR
         LOGICAL RETRY
         Logical flag RETRY may be set to .FALSE. to suppress repitition
  С
         of question after an incorrect response. In this case, an
  С
         error message is printed and execution is terminated. This feature
  С
         may be useful in the WWW version of the code, which is not truly
  С
         interactive.
  C
         DATA RETRY/.TRUE./
  С
         IF (IERR.NE.O) THEN
              IF (.NOT.RETRY) THEN
                   WRITE(6,667)
                   FORMAT('@ 00001 ABNORMAL TERMINATION: ',
     667
                   /,1x,' ERROR IN RETRY_INPUT: ',
        &
                   /,lx,' ERROR in user-supplied input. STOP')
                   STOP
              ELSE
                   WRITE(6,665) IERR
                   FORMAT(/,' ERROR ON INPUT: VAX ERROR CODE = ', I5,
     665
                        ' PLEASE TRY AGAIN.')
        &
                   IERR=0
             ENDIF
          ENDIF
          RETURN
          END
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for element IZ, energy EN (MeV/nuc) in one of two modes:
    IMODE=1: "Worst Day": particle fluxes based on observations from
              GOES (protons) and IMP-8 (heavy ions) for the 18-hour
              period beginning at 1300 UT on 20 OCT 1989.
    IMODE=2: "Worst Week": particle fluxes based on observations from
              GOES (protons) and IMP-8 (heavy ions) for the 180-hour
              period beginning at 1300 UT on 19 OCT 1989.
              NOTE: The actual termination time here is somewhat arbitrary,
                      since only a few percent of the flux was accumulated
    IMODE=3: Peak flux, based on peak 5-minute average flux observed on
     Average particle flux for this specified period is returned in
       REAL*4 EN, HOURS, SOLAR PROTONS, ERRFLUX, SOLAR HEAVY_IONS
       DIMENSION HOURS (3), NSEP (2), LSEP (4)
       DATA HOURS/18.0,180.0,0.083333/
       DATA NSEP/1,4/
       DATA LSEP/'200CT89','190CT89','220CT89','240CT89'/
       SEP FLUX=0.0
       IF (EN.LT.1.0 .or. EN.GT. 1.0E+5) RETURN
     IF (IMODE.LT.1 .or. IMODE.GT.3) RETURN
       IF (IZ.EQ.1) THEN
          ETEMP=EN
          IF (ETEMP.GT.600.) ETEMP=600.
          IF (IMODE.LT.3) THEN
          DO 1000 K=1, NSEP (IMODE)
            LABEL=LSEP(K)
            SEP_FLUX=SEP_FLUX+SOLAR_PROTONS(ETEMP,LABEL,ERRFLUX)
          CONTINUE
1000
          ELSEIF (IMODE.EQ.3) THEN
            LABEL='PEAKFLX'
             SEP FLUX=SOLAR_PROTONS(ETEMP, LABEL, ERRFLUX)
          Match high-energy extrapolation to alpha spectrum; AJT 12/27/96
C
           IF (IMODE.NE.2) SEP_FLUX=SEP_FLUX*(ETEMP/EN)**4.14060
           IF (IMODE.EQ.2) SEP_FLUX=SEP_FLUX*(ETEMP/EN)**3.76100
        ELSEIF (IZ.EQ.2) THEN
            IUSE=6
            Use D. Reames nominal He/C ratio:
C
            SEP_FLUX=122.5*SOLAR_HEAVY_IONS(IUSE,EN,IMODE,ERRFLUX)
            Fine-tune using the GOES alpha flux at ~7.3 MeV/nuc
C
            IF (IMODE.NE.2) SEP_FLUX=0.85*SEP_FLUX
```

```
C SHELLIG.FOR, Version 2.0, January 1992
   C
   C 11/1/91 SHELLG: lowest starting point for B0 search is 2
   C 1/27/92 Adopted to IGRF-91 coeffcients model
   C 2/5/92 Reduce variable-names: INTER(P)SHC,EXTRA(P)SHC,INITI(ALI)ZE
   C 12/9/92 DGRF/IGRF file names changed by AJT
            Changes in FELDCOF, for initialization purposes
   SUBROUTINES FINDBO, SHELLG, STOER, FELDG, FELDCOF, GETSHC.
          INTERSHC, EXTRASHC, INITIZE
   SUBROUTINE FINDBO (STPS, BDEL, VALUE, BEQU, RRO)
   C-----
   C FINDS SMALLEST MAGNETIC FIELD STRENGTH ON FIELD LINE
  C INPUT:
           STPS STEP SIZE FOR FIELD LINE TRACING
   C
          COMMON/FIDBO/
  C
            SP
                  DIPOLE ORIENTED COORDINATES FORM SHELLG; P(1,*),
                  P(2,*), P(3,*) CLOSEST TO MAGNETIC EQUATOR
  C
            BDEL
                  REQUIRED ACCURACY = [ B(LAST) - BEQU ] / BEQU
  C
                  B(LAST) IS FIELD STRENGTH BEFORE BEQU
  С
  C OUTPUT: VALUE =.FALSE., IF BEQU IS NOT MINIMAL VALUE ON FIELD LINE
  C
            BEQU MAGNETIC FIELD STRENGTH AT MAGNETIC EQUATOR
                  EQUATORIAL RADIUS NORMALIZED TO EARTH RADIUS
  C
            RRO
            BDEL FINAL ACHIEVED ACCURACY
u
        DIMENSION
                       P(8,4),SP(3)
LOGICAL
                      VALUE
Ŧ
        COMMON/FIDBO/
                       SP
STEP=STPS
L
        IRUN=0
7777 IRUN=IRUN+1
Ū
        IF (IRUN.GT.5) THEN
         VALUE=.FALSE.
         GOTO 8888
         ENDIF
  P(1,2) = SP(1)
        P(2,2) = SP(2)
        P(3,2) = SP(3)
        STEP=-SIGN(STEP, P(3,2))
        CALL STOER (P(1,2), BQ2, R2)
        P(1,3)=P(1,2)+0.5*STEP*P(4,2)
        P(2,3) = P(2,2) + 0.5*STEP*P(5,2)
        P(3,3) = P(3,2) + 0.5 * STEP
        CALL STOER (P(1,3), BQ3, R3)
        P(1,1) = P(1,2) - STEP*(2.*P(4,2) - P(4,3))
        P(2,1)=P(2,2)-STEP*(2.*P(5,2)-P(5,3))
        P(3,1) = P(3,2) - STEP
        CALL STOER (P(1,1), BQ1, R1)
        P(1,3)=P(1,2)+STEP*(20.*P(4,3)-3.*P(4,2)+P(4,1))/18.
        P(2,3) = P(2,2) + STEP*(20.*P(5,3) - 3.*P(5,2) + P(5,1)) / 18.
       P(3,3) = P(3,2) + STEP
       CALL STOER (P(1,3), BQ3, R3)
```

```
IF(BQ3.LE.BQ1) GOTO 2
           STEP=-STEP
           R3=R1
           BQ3=BQ1
           DO 1 I=1,5
                    ZZ=P(I,1)
                   P(I,1) = P(I,3)
   1
                   P(I,3)=ZZ
   C******************INITIALIZATION
           STEP12=STEP/12.
           VALUE=.TRUE.
           BMIN=1.E4
           BOLD=1.E4
   5555 P(1,3) = P(1,2) + STEP12 * (5.*P(4,3) + 8.*P(4,2) - P(4,1))
           N=N+1
         P(2,3) = P(2,2) + STEP12*(5.*P(5,3) + 8.*P(5,2) - P(5,1))
   C*******************PREDICTOR (FIELD LINE TRACING)
         P(1,4) = P(1,3) + STEP12*(23.*P(4,3)-16.*P(4,2)+5.*P(4,1))
         P(2,4) = P(2,3) + STEP12*(23.*P(5,3)-16.*P(5,2)+5.*P(5,1))
         P(3,4) = P(3,3) + STEP
         CALL STOER (P(1,4), BQ3,R3)
           DO 1111 J=1,3
           DO 1111 I=1,8
   1111
           P(I,J) = P(I,J+1)
           B=SQRT (BQ3)
           IF(B.LT.BMIN) BMIN=B
IF (B.LE.BOLD) THEN
                   BOLD=B
Lij
                   ROLD=1./R3
garde
garde
                   SP(1) = P(1,4)
Œ.
                   SP(2) = P(2,4)
SP(3) = P(3,4)
FL
                   GOTO 5555
                   ENDIF
i d
           IF (BOLD.NE.BMIN) THEN
E.
                   VALUE=.FALSE.
                   ENDIF
           BDELTA= (B-BOLD) /BOLD
           IF (BDELTA.GT.BDEL) THEN
                   STEP=STEP/10.
                   GOTO 7777
                   ENDIF
   8888
           RR0=ROLD
           BEQU=BOLD
           BDEL=BDELTA
           RETURN
           END
   C
         SUBROUTINE SHELLG (GLAT, GLON, ALT, DIMO, FL, ICODE, BO)
   C-----
   C CALCULATES L-VALUE FOR SPECIFIED GEODAETIC COORDINATES, ALTITUDE
  C AND GEMAGNETIC FIELD MODEL.
  C REF: G. KLUGE, EUROPEAN SPACE OPERATIONS CENTER, INTERNAL NOTE
  C
         NO. 67, 1970.
  С
         G. KLUGE, COMPUTER PHYSICS COMMUNICATIONS 3, 31-35, 1972
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C CHANGES (D. BILITZA, NOV 87):
     - USING CORRECT DIPOL MOMENT I.E., DIFFERENT COMMON/MODEL/
       - USING IGRF EARTH MAGNETIC FIELD MODELS FROM 1945 TO 1990
   C-----
   C INPUT: ENTRY POINT SHELLG
                  GLAT GEODETIC LATITUDE IN DEGREES (NORTH)
   C
   C
                  GLON GEODETIC LONGITUDE IN DEGREES (EAST)
   С
                  ALT ALTITUDE IN KM ABOVE SEA LEVEL
   С
             ENTRY POINT SHELLC
   C
                 V(3) CARTESIAN COORDINATES IN EARTH RADII (6371.2 KM)
   С
   C
                         X-AXIS POINTING TO EQUATOR AT 0 LONGITUDE
                         Y-AXIS POINTING TO EQUATOR AT 90 LONG.
   C
                         Z-AXIS POINTING TO NORTH POLE
   С
   C
                      DIPOL MOMENT IN GAUSS (NORMALIZED TO EARTH RADIUS)
   С
             DIMO
   С
   С
             COMMON
   С
                 X(3) NOT USED
   C
                 H(144) FIELD MODEL COEFFICIENTS ADJUSTED FOR SHELLG
   C-----
   C OUTPUT: FL
                       L-VALUE
   C
            ICODE
                       =1 NORMAL COMPLETION
                         =2 UNPHYSICAL CONJUGATE POINT (FL MEANINGLESS)
   C
                         =3 SHELL PARAMETER GREATER THAN LIMIT UP TO
  C
  C
                           WHICH ACCURATE CALCULATION IS REQUIRED;
                           APPROXIMATION IS USED.
             BO MAGNETIC FIELD STRENGTH IN GAUSS
  C-----
       DIMENSION V(3), U(3,3), P(8,100), SP(3)
L] C
       The following was an unlabeled common, which I have appropriately
       named. AJT 12-30-92.
        COMMON/BASTARDS/
                                X(3), H(144)
        COMMON/FIDBO/ SP
        COMMON/GENER/ UMR, ERA, AQUAD, BQUAD
   C-- RMIN, RMAX ARE BOUNDARIES FOR IDENTIFICATION OF ICODE=2 AND 3
   C-- STEP IS STEP SIZE FOR FIELD LINE TRACING
   C-- STEQ IS STEP SIZE FOR INTEGRATION
   С
        DATA RMIN, RMAX /0.05, 1.01/
        DATA STEP, STEQ /0.20, 0.03/
          BEQU=1.E10
   C****ENTRY POINT SHELLG TO BE USED WITH GEODETIC CO-ORDINATES
                                                                    SHEL0080
        RLAT=GLAT*UMR
        CT=SIN(RLAT)
                                                                    SHEL0100
        ST=COS(RLAT)
                                                                    SHEL0110
        D=SQRT (AQUAD- (AQUAD-BQUAD) *CT*CT)
        X(1) = (ALT + AQUAD/D) *ST/ERA
        X(3) = (ALT + BOUAD/D) *CT/ERA
        RLON=GLON*UMR
        X(2) = X(1) *SIN(RLON)
                                                                    SHEL0160
        X(1) = X(1) * COS(RLON)
                                                                    SHEL0170
        GOTO9
                                                                    SHEL0180
        ENTRY SHELLC (V, FL, BO)
                                                                    SHEL0190
   C*****ENTRY POINT SHELLC TO BE USED WITH CARTESIAN CO-ORDINATES
                                                                    SHEL0200
        X(1) = V(1)
                                                                    SHEL0210
        X(2) = V(2)
                                                                    SHEL0220
        X(3) = V(3)
                                                                    SHEL0230
   C*****CONVERT TO DIPOL-ORIENTED CO-ORDINATES
                                                                    SHEL0240
```

```
Α
                                    +0.9335804,+0.3583680,+0.0000000,
                                                                                  SHEL0260
                                    +0.0714471,-0.1861260,+0.9799247/
         B
                                                                                  SHEL0270
    9
          RQ=1./(X(1)*X(1)+X(2)*X(2)+X(3)*X(3))
          R3H=SQRT (RQ*SQRT (RQ))
                                                                                  SHEL0290
          P(1,2) = (X(1)*U(1,1)+X(2)*U(2,1)+X(3)*U(3,1))*R3H
                                                                                  SHEL0300
          P(2,2) = (X(1)*U(1,2)+X(2)*U(2,2)
                                                         ) *R3H
                                                                                  SHEL0310
          P(3,2) = (X(1)*U(1,3)+X(2)*U(2,3)+X(3)*U(3,3))*RQ
                                                                                  SHEL0320
    C****FIRST THREE POINTS OF FIELD LINE
                                                                                  SHEL0330
          STEP=-SIGN(STEP, P(3,2))
                                                                                  SHEL0340
          CALL STOER (P(1,2), BQ2, R2)
                                                                                  SHEL0350
          B0=SQRT (BQ2)
                                                                                  SHEL0360
          P(1,3) = P(1,2) + 0.5 * STEP * P(4,2)
                                                                                  SHEL0370
          P(2,3) = P(2,2) + 0.5 * STEP * P(5,2)
                                                                                  SHEL0380
          P(3,3)=P(3,2)+0.5*STEP
                                                                                  SHEL0390
          CALL STOER (P(1,3), BQ3, R3)
                                                                                  SHEL0400
          P(1,1) = P(1,2) - STEP*(2.*P(4,2) - P(4,3))
                                                                                  SHEL0410
          P(2,1)=P(2,2)-STEP*(2.*P(5,2)-P(5,3))
                                                                                  SHEL0420
          P(3,1) = P(3,2) - STEP
                                                                                  SHEL0430
          CALL STOER (P(1,1), BQ1, R1)
                                                                                  SHEL0440
          P(1,3)=P(1,2)+STEP*(20.*P(4,3)-3.*P(4,2)+P(4,1))/18.
                                                                                  SHEL0450
          P(2,3) = P(2,2) + STEP*(20.*P(5,3)-3.*P(5,2)+P(5,1))/18.
                                                                                  SHEL0460
          P(3,3) = P(3,2) + STEP
                                                                                  SHEL0470
          CALL STOER (P(1,3), BQ3, R3)
                                                                                  SHEL0480
   C*****INVERT SENSE IF REQUIRED
                                                                                  SHEL0490
Ī
          IF (BO3.LE.BO1) GOTO2
                                                                                  SHEL0500
          STEP=-STEP
                                                                                  SHEL0510
          R3=R1
                                                                                  SHEL0520
Ŧij.
          BQ3=BQ1
                                                                                 SHEL0530
          DO 1 I=1,7
                                                                                 SHEL0540
          ZZ=P(I,1)
                                                                                 SHEL0550
          P(I,1) = P(I,3)
                                                                                 SHEL0560
          P(I,3)=ZZ
                                                                                 SHEL0570
   C****SEARCH FOR LOWEST MAGNETIC FIELD STRENGTH
1
          IF (BQ1.LT.BEQU) THEN
FL
            BEQU=BQ1
IJ
            IEOU=1
ENDIF
          IF (BQ2.LT.BEQU) THEN
            BEQU=BQ2
            IEQU=2
            ENDIF
          IF (BO3.LT.BEOU) THEN
            BEQU=BQ3
            IEOU=3
            ENDIF
   C*****INITIALIZATION OF INTEGRATION LOOPS
                                                                                 SHEL0580
          STEP12=STEP/12.
          STEP2=STEP+STEP
                                                                                 SHEL0600
          STEQ=SIGN(STEQ, STEP)
                                                                                 SHEL0610
          FT=0.
                                                                                 SHEL0620
          ICODE=1
                                                                                 SHEL0630
          ORADIK=0.
                                                                                 SHEL0640
          OTERM=0.
                                                                                 SHEL0650
          STP=R2*STEO
                                                                                 SHEL0660
          Z=P(3,2)+STP
                                                                                 SHEL0670
         STP=STP/0.75
         P(8,1) = STEP2*(P(1,1)*P(4,1)+P(2,1)*P(5,1))
                                                                                 SHEL0690
         P(8,2) = STEP2*(P(1,2)*P(4,2)+P(2,2)*P(5,2))
                                                                                 SHEL0700
   C****MAIN LOOP (FIELD LINE TRACING)
                                                                                 SHEL0710
```

+0.3511737,-0.9148385,-0.1993679,

SHEL0250

DATA U/

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C****PREPARE EXPANSION COEFFICIENTS FOR INTERPOLATION
    C****OF SLOWLY VARYING QUANTITIES
           P(8,N) = STEP2*(P(1,N)*P(4,N)+P(2,N)*P(5,N))
           C0=P(1,N-1)**2+P(2,N-1)**2
           C1=P(8,N-1)
           C2 = (P(8,N) - P(8,N-2)) *0.25
           C3 = (P(8,N) + P(8,N-2) - C1 - C1) / 6.0
           D0=P(6,N-1)
           D1 = (P(6, N) - P(6, N-2)) *0.5
           D2 = (P(6, N) + P(6, N-2) - D0 - D0) * 0.5
           E0=P(7,N-1)
           E1=(P(7,N)-P(7,N-2))*0.5
           E2 = (P(7,N) + P(7,N-2) - E0 - E0) * 0.5
    C*****INNER LOOP (FOR QUADRATURE)
           T = (Z - P(3, N-1)) / STEP
           IF (T.GT.1.) GOTO5
           HLI=0.5*(((C3*T+C2)*T+C1)*T+C0)
           R=HLI+SQRT (HLI*HLI+ZQ)
           IF (R.LE.RMIN) GOTO30
          RO=R*R
           FF=SQRT(1.+3.*ZQ/RQ)
           RADIK=B0-((D2*T+D1)*T+D0)*R*RQ*FF
           IF (R-RMAX) 44, 44, 45
    45
           ICODE=2
          RADIK=RADIK-12.*(R-RMAX)**2
           IF(RADIK+RADIK.LE.ORADIK) GOTO 10
           TERM=SQRT(RADIK)*FF*((E2*T+E1)*T+E0)/(RQ+ZQ)
ļ.
           FI=FI+STP* (OTERM+TERM)
*
          ORADIK=RADIK
          OTERM=TERM
          STP=R*STEO
          Z=Z+STP
.
Pratic
          GOTO4
   C****PREDICTOR (FIELD LINE TRACING)
          P(1, N+1) = P(1, N) + STEP12 * (23.*P(4, N) - 16.*P(4, N-1) + 5.*P(4, N-2))
           P(2,N+1) = P(2,N) + STEP12 * (23.*P(5,N)-16.*P(5,N-1)+5.*P(5,N-2))
          P(3,N+1) = P(3,N) + STEP
          CALL STOER (P(1,N+1),BQ3,R3)
    C*****SEARCH FOR LOWEST MAGNETIC FIELD STRENGTH
           IF (BQ3.LT.BEQU) THEN
             IEQU=N+1
            BEOU=BO3
            ENDIF
          CONTINUE
    3
    10
          IF(IEQU.lt.2) IEQU=2
          SP(1) = P(1, IEQU-1)
          SP(2) = P(2, IEQU-1)
          SP(3) = P(3, IEQU-1)
          IF (ORADIK.LT.1E-15) GOTO11
          FI=FI+STP/0.75*OTERM*ORADIK/(ORADIK-RADIK)
```

C-- The minimal allowable value of FI was changed from 1E-15 to 1E-12, C-- because 1E-38 is the minimal allowable arg. for ALOG in our envir.

C-- D. Bilitza, Nov 87.

C

P(1,N) = P(1,N-1) + STEP12*(5.*P(4,N)+8.*P(4,N-1)-P(4,N-2))

P(2,N) = P(2,N-1) + STEP12 * (5.*P(5,N) + 8.*P(5,N-1) - P(5,N-2))

SHEL0720

SHEL0730

SHEL0740

SHEL0750

SHEL0760

SHEL0770

SHEL0780

SHEL-0790

SHEL0800

SHEL0810

SHEL0830

SHEL0840

SHEL0850

SHEL0870

SHEL0880

SHEL0890

SHEL0900

SHEL0910

SHEL0920

SHEL0950

SHEL0970

SHEL0980

SHEL0990

SHEL1000

SHEL1010

SHEL1030

SHEL1040

SHEL1050

SHEL1060

SHEL1070

SHEL1080

SHEL1090

SHEL1100

SHEL1110

SHEL1120

SHEL1130

SHEL1140

SHEL1150

DO 3 N=3,3333

C*****CORRECTOR (FIELD LINE TRACING)

```
FI=0.5*ABS(FI)/SQRT(B0)+1E-12
C*****COMPUTE L FROM B AND I. SAME AS CARMEL IN INVAR.
C-- Correct dipole moment is used here. D. Bilitza, Nov 87.
C
     DIMOB0=DIMO/B0
     XX=ALOG(FI*FI*FI/DIMOB0+1E-12) !added 1E-12, 5-14-96, PRB.
      IF(XX.GT.23.0) GOTO 776
     IF(XX.GT.11.7) GOTO 775
     IF(XX.GT.+3.0) GOTO 774
     IF(XX.GT.-3.0) GOTO 773
     IF(XX.GT.-22.) GOTO 772
  771 GG=3.33338E-1*XX+3.0062102E-1
                                                                      SHEL1250
     GOTO777
                                                                      SHEL1260
  772 GG=(((((((-8.1537735E-14*XX+8.3232531E-13)*XX+1.0066362E-9)*XX+ SHEL1270
    18.1048663E-8) *XX+3.2916354E-6) *XX+8.2711096E-5) *XX+1.3714667E-3) * SHEL1280
    2XX+1.5017245E-2) *XX+4.3432642E-1) *XX+6.2337691E-1
                                                                     SHEL1290
     GOTO777
                                                                     SHEL1300
  773 GG=((((((((2.6047023E-10*XX+2.3028767E-9)*XX-2.1997983E-8)*XX-
                                                                     SHEL1310
    15.3977642E-7) *XX-3.3408822E-6) *XX+3.8379917E-5) *XX+1.1784234E-3) * SHEL1320
    2XX+1.4492441E-2)*XX+4.3352788E-1)*XX+6.228644E-1
                                                                     SHEL1330
                                                                     SHEL1340
  774 GG=(((((((6.3271665E-10*XX-3.958306E-8)*XX+9.9766148E-07)*XX-
                                                                     SHEL1350
    11.2531932E-5) *XX+7.9451313E-5) *XX-3.2077032E-4) *XX+2.1680398E-3) * SHEL1360
    2XX+1.2817956E-2) *XX+4.3510529E-1) *XX+6.222355E-1
                                                                     SHEL1370
     GOTO777
                                                                     SHEL1 380
  775 GG=(((((2.8212095E-8*XX-3.8049276E-6)*XX+2.170224E-4)*XX-6.7310339SHEL1390
    1E-3) *XX+1.2038224E-1) *XX-1.8461796E-1) *XX+2.0007187E0
                                                                     SHEL1400
     GOTO777
                                                                     SHEL1410
  776 GG=XX-3.0460681E0
                                                                     SHEL1420
  777 FL=EXP(ALOG((1.+EXP(GG))*DIMOB0)/3.0)
                                                                     SHEL1440
C****APPROXIMATION FOR HIGH VALUES OF L.
                                                                     SHEL1450
     ICODE=3
                                                                     SHEL1460
     T=-P(3,N-1)/STEP
                                                                     SHEL1470
     FL=1./(ABS(((C3*T+C2)*T+C1)*T+C0)+1E-15)
                                                                     SHEL1480
     RETURN
                                                                     SHEL1490
     END
                                                                     SHET-1500
C
C
     SUBROUTINE STOER (P, BQ, R)
                                                                     SHEL1510
C* SUBROUTINE USED FOR FIELD LINE TRACING IN SHELLG
C* CALLS ENTRY POINT FELDI IN GEOMAGNETIC FIELD SUBROUTINE FELDG
DIMENSION
                      P(7), U(3,3)
C
     The following was an unlabeled common, which I have appropriately
C
     named. AJT 12-30-92.
     COMMON/BASTARDS/
                              XI(3),H(144)
C****XM,YM,ZM ARE GEOMAGNETIC CARTESIAN INVERSE CO-ORDINATES
                                                                     SHEL1540
     ZM=P(3)
                                                                     SHEL1550
     FLI=P(1)*P(1)+P(2)*P(2)+1E-15
     R=0.5*(FLI+SQRT(FLI*FLI+(ZM+ZM)**2))
     RQ=R*R
     WR=SQRT(R)
                                                                     SHEL1590
     XM=P(1)*WR
                                                                     SHEL1600
     YM=P(2)*WR
                                                                     SHEL1610
C****TRANSFORM TO GEOGRAPHIC CO-ORDINATE SYSTEM
                                                                     SHEL1620
     DATA U/
                            +0.3511737, -0.9148385, -0.1993679,
                                                                     SHEL1630
    Α
                            +0.9335804,+0.3583680,+0.0000000,
                                                                     SHEL1640
```

```
+0.0714471,-0.1861260,+0.9799247/
                                                                         SHEL:1650
     В
      XI(1) = XM*U(1,1) + YM*U(1,2) + ZM*U(1,3)
                                                                         SHEL1660
                                                                         SHEL1670
      XI(2) = XM*U(2,1) + YM*U(2,2) + ZM*U(2,3)
      XI(3) = XM*U(3,1)
                                                                         SHEL1680
C*****COMPUTE DERIVATIVES
                                                                         SHEL1690
                                                                          SHEL1700
       CALL FELDI(XI, H)
                                                                         SHEL1700
      CALL FELDI
                                                                         SHEL1710
      Q=H(1)/RQ
                                                                         SHEL1720
      DX=H(3)+H(3)+Q*XI(1)
                                                                         SHEL1730
      DY=H(4)+H(4)+Q*XI(2)
                                                                         SHEL1740
      DZ=H(2)+H(2)+O*XI(3)
C****TRANSFORM BACK TO GEOMAGNETIC CO-ORDINATE SYSTEM
                                                                         SHEL-1750
                                                                         SHEL1760
      DXM=U(1,1)*DX+U(2,1)*DY+U(3,1)*DZ
                                                                         SHEL1770
      DYM=U(1,2)*DX+U(2,2)*DY
                                                                         SHEL1780
      DZM=U(1,3)*DX+U(2,3)*DY+U(3,3)*DZ
      DR = (XM*DXM+YM*DYM+ZM*DZM) / R
                                                                         SHEL1790
C*****FORM SLOWLY VARYING EXPRESSIONS
                                                                         SHEL1800
      P(4) = (WR*DXM-0.5*P(1)*DR) / (R*DZM)
                                                                         SHEL1810
      P(5) = (WR*DYM-0.5*P(2)*DR) / (R*DZM)
                                                                         SHEL1820
      DSO=RQ* (DXM*DXM+DYM*DYM+DZM*DZM)
      BQ=DSQ*RQ*RQ
      P(6) = SQRT(DSQ/(RQ+3.*ZM*ZM))
                                                                         SHEL1850
      P(7) = P(6) * (RQ + ZM * ZM) / (RQ * DZM)
                                                                         SHEL1860
                                                                         SHEL1870
      RETURN
                                                                         SHEL1880
      END
С
                                                                         SHEL1890
      SUBROUTINE FELDG (GLAT, GLON, ALT, BNORTH, BEAST, BDOWN, BABS)
C-----
C CALCULATES EARTH MAGNETIC FIELD FROM SPHERICAL HARMONICS MODEL
C REF: G. KLUGE, EUROPEAN SPACE OPERATIONS CENTRE, INTERNAL NOTE 61,
       1970.
C CHANGES (D. BILITZA, NOV 87):
    - FIELD COEFFICIENTS IN BINARY DATA FILES INSTEAD OF BLOCK DATA
    - CALCULATES DIPOL MOMENT
C INPUT: ENTRY POINT FELDG
                GLAT GEODETIC LATITUDE IN DEGREES (NORTH)
С
С
                GLON GEODETIC LONGITUDE IN DEGREES (EAST)
C
                ALT
                     ALTITUDE IN KM ABOVE SEA LEVEL
C
           ENTRY POINT FELDC
C
C
                V(3) CARTESIAN COORDINATES IN EARTH RADII (6371.2 KM)
C
                        X-AXIS POINTING TO EQUATOR AT 0 LONGITUDE
C
                        Y-AXIS POINTING TO EQUATOR AT 90 LONG.
C
                        Z-AXIS POINTING TO NORTH POLE
C
           COMMON BLANK AND ENTRY POINT FELDI ARE NEEDED WHEN USED
C
C
             IN CONNECTION WITH L-CALCULATION PROGRAM SHELLG.
C
C
           COMMON /MODEL/ AND /GENER/
                        = ATAN(1.0)*4./180. <DEGREE>*UMR=<RADIANT>
C
                TIMR
C
                ERA
                        EARTH RADIUS FOR NORMALIZATION OF CARTESIAN
                        COORDINATES (6371.2 KM)
C
C
                AQUAD, BQUAD
                              SOUARE OF MAJOR AND MINOR HALF AXIS FOR
C
                        EARTH ELLIPSOID AS RECOMMENDED BY INTERNATIONAL
C
                        ASTRONOMICAL UNION (6378.160, 6356.775 KM).
C
                        MAXIMUM ORDER OF SPHERICAL HARMONICS
                NMAX
                TIME
                        YEAR (DECIMAL: 1973.5) FOR WHICH MAGNETIC
C
```

C C	G (M)	FIELD IS TO BE CALCULATED NORMALIZED FIELD COEFFICIENTS (SEE FELDCOF) M=NMAX*(NMAX+2)		
C C C C C	BNORTH, BEAS TO TH POINT	TIC FIELD STRENGTH IN GAUSS T, BDOWN COMPONENTS OF THE FIELD WITH RESPECT E LOCAL GEODETIC COORDINATE SYSTEM, WITH AXIS ING IN THE TANGENTIAL PLANE TO THE NORTH, EAST OWNWARD.		
C	DIMENSION	V(3),B(3)		
	CHARACTER*30	. 3		
C	_	an unlabeled common, which I have appropriately	Y	
С	named. AJT 12-30	-92. XI(3),H(144)		
		NAME, NMAX, TIME, G(144)		
	COMMON/GENER/	UMR, ERA, AQUAD, BQUAD		
С				
C IS RECORDS ENTRY POINT C				
	IS=1	G TO BE USED WITH GEODETIC CO-ORDINATES	SHEL1920 SHEL1930	
	RLAT=GLAT*UMR CT=SIN(RLAT)		CUDI 1050	
	ST=COS (RLAT)		SHEL1950 SHEL1960	
	D=SQRT (AQUAD- (AQU	AD-BQUAD) *CT*CT)	SHEL1970	
	RLON=GLON*UMR			
	CP=COS (RLON)		SHEL1990	
	SP=SIN(RLON) ZZZ=(ALT+BQUAD/D RHO=(ALT+AQUAD/D		SHEL2000	
	XXX=RHO*CP		SHEL2030	
-	YYY=RHO*SP		SHEL2040	
	GOTO10		SHEL2050	
C***	ENTRY FELDC(V,B) **ENTRY POINT FELD	C TO BE USED WITH CARTESIAN CO-ORDINATES	SHEL2060 SHEL2070	
•	IS=2	o lo de oble with disciplination of old intilled	SHEL2090	
	XXX=V(1)		SHEL2100	
	YYY=V(2)		SHEL2110	
10	ZZZ=V(3)	V+VVV , PPP+PPP \	SHEL2120	
10	RQ=1./(XXX*XXX+YY) XI(1)=XXX*RQ	Y ~ Y Y Y + Z Z Z ~ Z Z Z)	SHEL2140	
	XI(2)=YYY*RQ		SHEL2150	
	XI(3) = ZZZ*RQ		SHEL2160	
	GOTO20		SHEL2170	
G + + +	ENTRY FELDI	T VIGED FOR I GOVERNMENT	SHEL2180	
CAAA	**ENTRY POINT FELD. IS=3	I USED FOR L COMPUTATION	SHEL2190 SHEL2200	
20	IHMAX=NMAX*NMAX+1		SHEL2210	
	LAST=IHMAX+NMAX+N	XAM	SHEL2220	
	IMAX=NMAX+NMAX-1		SHEL2230	
	DO 8 I=IHMAX, LAST		SHEL2240	
8	H(I) = G(I) DO 6 K=1,3,2		SHEL2250	
	I=IMAX		SHEL2260 SHEL2270	
	IH=IHMAX		SHEL2280	
1	IL=IH-I		SHEL2290	
	F=2./FLOAT(I-K+2)		SHEL2300	
	X=XI(1)*F Y=XI(2)*F		SHEL2310	
	I=VT /7/ "L		SHEL2320	

```
Z=XI(3)*(F+F)
                                                                           SHEL2330
       I=I-2
                                                                           SHEL2340
       IF(I-1)5,4,2
                                                                           SHEL2350
2
       DO 3 M=3, I, 2
                                                                           SHEL2360
       H(IL+M+1) = G(IL+M+1) + Z*H(IH+M+1) + X*(H(IH+M+3) - H(IH+M-1))
                                                                           SHEL2370
                                      -Y*(H(IH+M+2)+H(IH+M-2))
                                                                           SHEL2380
3
      H(IL+M) = G(IL+M) + Z+H(IH+M) + X+(H(IH+M+2) - H(IH+M-2))
                                                                           SHEL2390
                                +Y*(H(IH+M+3)+H(IH+M-1))
                                                                           SHEL2400
4
      H(IL+2) = G(IL+2) + Z*H(IH+2) + X*H(IH+4) - Y*(H(IH+3) + H(IH))
                                                                           SHEL2410
      H(IL+1) = G(IL+1) + Z*H(IH+1) + Y*H(IH+4) + X*(H(IH+3) - H(IH))
                                                                           SHEL2420
5
      H(IL) = G(IL) + Z*H(IH) + 2.*(X*H(IH+1) + Y*H(IH+2))
                                                                           SHEL2430
       IH=IL
                                                                           SHEL2440
       IF (I.GE.K) GOTO1
                                                                           SHEL2450
6
      CONTINUE
                                                                           SHEL2460
       IF (IS.EO.3) RETURN
                                                                           SHEL2470
       S=.5*H(1)+2.*(H(2)*XI(3)+H(3)*XI(1)+H(4)*XI(2))
                                                                           SHEL2480
       T = (RQ + RQ) * SQRT (RQ)
                                                                           SHEL2490
      BXXX=T*(H(3)-S*XXX)
                                                                           SHEL2500
      BYYY=T*(H(4)-S*YYY)
                                                                           SHEL2510
      BZZZ=T*(H(2)-S*ZZZ)
                                                                           SHEL2520
      IF(IS.EQ.2)GOTO7
                                                                           SHEL2530
      BABS=SQRT (BXXX*BXXX+BYYY*BYYY+BZZZ*BZZZ)
      BEAST=BYYY*CP-BXXX*SP
                                                                          SHEL2550
      BRHO=BYYY*SP+BXXX*CP
                                                                          SHEL2560
      BNORTH=BZZZ*ST-BRHO*CT
                                                                          SHEL2570
      BDOWN=-BZZZ*CT-BRHO*ST
                                                                          SHEL2580
      RETURN
                                                                          SHEL2590
      B(1) = BXXX
                                                                          SHEL2600
      B(2) = BYYY
                                                                          SHEL2610
      B(3) = BZZZ
                                                                          SHEL2620
      RETURN
                                                                          SHEL2630
      END
                                                                          SHEL2640
С
        SUBROUTINE FELDCOF (YEAR, DIMO)
C DETERMINES COEFFICIENTS AND DIPOL MOMENT FROM IGRF MODELS
C
С
        INPUT: YEAR
                         DECIMAL YEAR FOR WHICH GEOMAGNETIC FIELD IS TO
C
                        BE CALCULATED
C
        OUTPUT: DIMO
                         GEOMAGNETIC DIPOL MOMENT IN GAUSS (NORMALIZED
C
                        TO EARTH'S RADIUS) AT THE TIME (YEAR)
C D. BILITZA, NSSDC, GSFC, CODE 633, GREENBELT, MD 20771.
C
        (301) 286-9536 NOV 1987.
C
C Modified by AJT 12-9-92:
C allow for multiple calls: field paramters are not read in unless
C year value is changed. Call to INITIZE also added here.
C-----
                      FILMOD, FIL1, FIL2 !5-14-96, change from 17 to 30
        CHARACTER*30
                        GH1 (144), GH2 (120), GHA (144), FILMOD (11), DTEMOD (11)
        DIMENSION
        DOUBLE PRECISION X, FO, F
        COMMON/MODEL/
                        FIL1, NMAX, TIME, GH1
        COMMON/GENER/
                        UMR, ERAD, AQUAD, BQUAD
        DATA
                        FILMOD/
C
      &
                         'creme96:dgrf45.dat',
С
                         'creme96:dgrf50.dat',
C
      1
                         'creme96:dgrf55.dat', 'creme96:dgrf60.dat',
C
      &
                         'creme96:dqrf65.dat',
```

```
C
                          'creme96:dgrf70.dat', 'creme96:dgrf75.dat',
       2
 C
                           'creme96:dgrf80.dat',
 C
       3
                          'creme96:dgrf85.dat', 'creme96:igrf90.dat',
 С
                           'creme96:igrf90s.dat'/
 С
         Remove directory path, per BB's new file open routines AJT 11/18/97
      &
                          'dgrf45.dat',
      £
                          'dgrf50.dat',
      1
                          'dgrf55.dat', 'dgrf60.dat',
                          'dgrf65.dat',
      &
      2
                          'dgrf70.dat', 'dgrf75.dat',
      &
                          'dgrf80.dat',
      3
                          'dgrf85.dat', 'igrf90.dat',
                          'igrf90s.dat'/
      æ
         DATA
                          DTEMOD / 1945., 1950., 1955., 1960., 1965.,
      1
                          1970., 1975., 1980., 1985., 1990., 1995./
         DATA YEAROLD/0/, IENT/0/
         COMMON/AJTDIMO/AJTDIMO
         IF (IENT.EQ.0) THEN
             IENT=1
             write(*,*)' Initialization call to FELDCOF: YEAR = ',YEAR
             CALL INITIZE
         ENDIF
         IF (ABS (YEAR-YEAROLD) .LT.0.001) THEN
             DIMO=AJTDIMO
             RETURN
        ENDIF
        YEAROLD=YEAR
   numye is number of years represented by IGRF models
C
        NUMYE=10
C
   IS=0 FOR SCHMIDT NORMALIZATION
                                     IS=1 GAUSS NORMALIZATION
   IU IS INPUT UNIT NUMBER FOR IGRF COEFFICIENT SETS
        IU = 10
        IS = 0
C-- DETERMINE IGRF-YEARS FOR INPUT-YEAR
        TIME = YEAR
        IYEA = INT(YEAR/5.)*5
        L = (IYEA - 1945)/5 + 1
        IF(L.LT.1) L=1
        IF (L.GT.NUMYE) L=NUMYE
        DTE1 = DTEMOD(L)
        FIL1 = FILMOD(L)
        DTE2 = DTEMOD(L+1)
        FIL2 = FILMOD(L+1)
C-- GET IGRF COEFFICIENTS FOR THE BOUNDARY YEARS
    Error messages added by AJT 11/24/97
        CALL GETSHC (IU, FIL1, NMAX1, ERAD, GH1, IER)
            IF (IER .NE. 0) THEN
                WRITE(6,9999) FIL1, IER
 9999
                FORMAT('@ 02001 ABNORMAL TERMINATION: ',
     &
               /,1x,' IGRF Coefficient file not found: ',
     &
               /,1x,A80,
                /,1x,' Error return code = ',I10,' STOP.')
                STOP
            ENDIF
```

```
CALL GETSHC (IU, FIL2, NMAX2, ERAD, GH2, IER)
            IF (IER .NE. 0) THEN
                WRITE(6,9999) FIL2, IER
                STOP
            ENDIF
C-- DETERMINE IGRF COEFFICIENTS FOR YEAR
        IF (L .LE. NUMYE-1) THEN
          CALL INTERSHC (YEAR, DTE1, NMAX1, GH1, DTE2,
     1
               NMAX2, GH2, NMAX, GHA)
        ELSE
          CALL EXTRASHC (YEAR, DTE1, NMAX1, GH1, NMAX2,
     1
               GH2, NMAX, GHA)
        ENDIF
C-- DETERMINE MAGNETIC DIPOL MOMENT AND COEFFIECIENTS G
        F0=0.D0
        DO 1234 J=1,3
          F = GHA(J) * 1.D-5
           FO = FO + F * F
1234
        CONTINUE
        DIMO = DSORT(F0)
        AJTDIMO=DIMO
        GH1(1) = 0.0
        I=2
        F0=1.D-5
        IF(IS.EQ.0) F0=-F0
        SQRT2=SQRT(2.)
     DO 9 N=1, NMAX
       X = N
       F0 = F0 * X * X / (4.D0 * X - 2.D0)
       IF(IS.EQ.0) F0 = F0 * (2.D0 * X - 1.D0) / X
       F = F0 * 0.5D0
       IF(IS.EQ.0) F = F * SQRT2
       GH1(I) = GHA(I-1) * F0
       I = I+1
     DO 9 M=1,N
       F = F * (X + M) / (X - M + 1.D0)
       IF(IS.EQ.0) F = F * DSQRT((X - M + 1.D0) / (X + M))
       GH1(I) = GHA(I-1) * F
       GH1(I+1) = GHA(I) * F
       I=I+2
9
      CONTINUE
       RETURN
       END
C
C
       SUBROUTINE GETSHC (IU, FSPEC, NMAX, ERAD, GH, IER)
C
С
       Version 1.01
C
С
       Reads spherical harmonic coefficients from the specified
C
       file into an array.
C
C
       Input:
С
                 - Logical unit number
С
           FSPEC - File specification
C
```

```
the true for the party party of the collection o
```

C

Output:

```
NMAX - Maximum degree and order of model
C
          ERAD - Earth's radius associated with the spherical
C
               harmonic coefficients, in the same units as
C
C
                elevation
               - Schmidt quasi-normal internal spherical
С
                harmonic coefficients
C
С
          IER - Error number: = 0, no error
                            = -2, records out of order
C
                            = FORTRAN run-time error number
С
C
C
      A. Zunde
      USGS, MS 964, Box 25046 Federal Center, Denver, CO 80225
C
CHARACTER
DIMENSION
                   FSPEC*(*)
                   GH(*)
       integer
                   stat,creme96_open
C -----
      Open coefficient file. Read past first header record.
      Read degree and order of model and Earth's radius.
C
C -----
    made READONLY, 5-16-96, PRB.
C
      OPEN (IU, FILE=FSPEC, STATUS='OLD', READONLY, IOSTAT=IER, ERR=999)
      stat = creme96 open(fspec,'cr96tables',iu,'old')
      if (stat .ne. 0) goto 999
      READ (IU, *, IOSTAT=IER, ERR=999)
       READ (IU, *, IOSTAT=IER, ERR=999) NMAX, ERAD
C -----
      Read the coefficient file, arranged as follows:
C
C
                                  ______
C
C
                                     0 GH(1) -
                                      1 GH(2) GH(3)
C
                                 1
                                        GH(4) -
                                 2
                                      0
C
                                 2 1 GH(5) GH(6)
С
                                2
                                     2 GH(7) GH(8)
С
          NMAX*(NMAX+3)/2
                                     0
C
             records
                                 3
                                          GH(9) -
С
С
C
          NMAX*(NMAX+2)
C
          elements in GH
                                NMAX NMAX
C
C
      N and M are, respectively, the degree and order of the
C
       coefficient.
       I = 0
       DO 2211 NN = 1, NMAX
          DO 2233 MM = 0, NN
             READ (IU, *, IOSTAT=IER, ERR=999) N, M, G, H
              IF (NN .NE. N .OR. MM .NE. M) THEN
                 IER = -2
                 GOTO 999
             ENDIF
```

```
I = I + 1
                GH(I) = G
                IF (M .NE. O) THEN
                    I = I + 1
                    GH(I) = H
                ENDIF
2233
            CONTINUE
2211
        CONTINUE
999
        CLOSE (IU)
        RETURN
        END
C
        SUBROUTINE INTERSHC (DATE, DTE1, NMAX1, GH1, DTE2,
                             NMAX2, GH2, NMAX, GH)
С
С
       Version 1.01
С
С
        Interpolates linearly, in time, between two spherical
С
       harmonic models.
C
С
       Input:
C
           DATE - Date of resulting model (in decimal year)
C
           DTE1 - Date of earlier model
C
           NMAX1 - Maximum degree and order of earlier model
C
                 - Schmidt quasi-normal internal spherical
С
                   harmonic coefficients of earlier model
С
           DTE2 - Date of later model
С
           NMAX2 - Maximum degree and order of later model
С
                - Schmidt quasi-normal internal spherical
С
                   harmonic coefficients of later model
С
C
       Output:
C
                 - Coefficients of resulting model
C
           NMAX - Maximum degree and order of resulting model
C
C
       A. Zunde
С
       USGS, MS 964, Box 25046 Federal Center, Denver, CO 80225
       DIMENSION
                       GH1(*), GH2(*), GH(*)
C
       The coefficients (GH) of the resulting model, at date
С
С
       DATE, are computed by linearly interpolating between the
С
       coefficients of the earlier model (GH1), at date DTE1,
C
       and those of the later model (GH2), at date DTE2. If one
С
       model is smaller than the other, the interpolation is
C
       performed with the missing coefficients assumed to be 0.
       FACTOR = (DATE - DTE1) / (DTE2 - DTE1)
       IF (NMAX1 .EQ. NMAX2) THEN
           K = NMAX1 * (NMAX1 + 2)
```

```
ELSE IF (NMAX1 .GT. NMAX2) THEN
               K = NMAX2 * (NMAX2 + 2)
               L = NMAX1 * (NMAX1 + 2)
               DO 1122 I = K + 1, L
                  GH(I) = GH1(I) + FACTOR * (-GH1(I))
   1122
               NMAX = NMAX1
           ELSE
               K = NMAX1 * (NMAX1 + 2)
               L = NMAX2 * (NMAX2 + 2)
               DO 1133 I = K + 1, L
                  GH(I) = FACTOR * GH2(I)
   1133
               NMAX = NMAX2
           ENDIF
           DO 1144 I = 1, K
               GH(I) = GH1(I) + FACTOR * (GH2(I) - GH1(I))
   1144
           RETURN
           END
   C
   C
           SUBROUTINE EXTRASHC (DATE, DTE1, NMAX1, GH1, NMAX2,
        1
                                GH2, NMAX, GH)
   C
C
           Version 1.01
   C
           Extrapolates linearly a spherical harmonic model with a
   С
           rate-of-change model.
   C
   С
   C
           Input:
Ξ
               DATE - Date of resulting model (in decimal year)
   С
               DTE1 - Date of base model
   C
C
               NMAX1 - Maximum degree and order of base model
                     - Schmidt quasi-normal internal spherical
   С
                      harmonic coefficients of base model
   C
               NMAX2 - Maximum degree and order of rate-of-change
   С
   C
                      model
   C
               GH2
                   - Schmidt quasi-normal internal spherical
   C
                      harmonic coefficients of rate-of-change model
   C
   С
           Output:
   C
                     - Coefficients of resulting model
   C
                    - Maximum degree and order of resulting model
   С
   C
           A. Zunde
   C
           USGS. MS 964, Box 25046 Federal Center, Denver, CO 80225
   C
                          GH1(*), GH2(*), GH(*)
           DIMENSION
    C -----
           The coefficients (GH) of the resulting model, at date
    C
    C
           DATE, are computed by linearly extrapolating the coef-
    С
           ficients of the base model (GH1), at date DTE1, using
    C
           those of the rate-of-change model (GH2), at date DTE2. If
```

one model is smaller than the other, the extrapolation is

C

NMAX = NMAX1

```
performed with the missing coefficients assumed to be 0.
       FACTOR = (DATE - DTE1)
        IF (NMAX1 .EQ. NMAX2) THEN
           K = NMAX1 * (NMAX1 + 2)
           NMAX = NMAX1
       ELSE IF (NMAX1 .GT. NMAX2) THEN
           K = NMAX2 * (NMAX2 + 2)
           L = NMAX1 * (NMAX1 + 2)
           DO 1155 I = K + 1, L
1155
               GH(I) = GH1(I)
           NMAX = NMAX1
       ELSE
           K = NMAX1 * (NMAX1 + 2)
           L = NMAX2 * (NMAX2 + 2)
           DO 1166 I = K + 1, L
               GH(I) = FACTOR * GH2(I)
1166
           NMAX = NMAX2
       ENDIF
       DO 1177 I = 1, K
1177
           GH(I) = GH1(I) + FACTOR * GH2(I)
       RETURN
       END
C
       SUBROUTINE INITIZE
C-----
C Initializes the parameters in COMMON/GENER/
C
C
               = ATAN(1.0)*4./180. <DEGREE>*UMR=<RADIANT>
       TIMR
C
       ERA
              EARTH RADIUS FOR NORMALIZATION OF CARTESIAN
С
                       COORDINATES (6371.2 KM)
C
      EREQU MAJOR HALF AXIS FOR EARTH ELLIPSOID (6378.160 KM)
C
       ERPOL MINOR HALF AXIS FOR EARTH ELLIPSOID (6356.775 KM)
C
       AQUAD SQUARE OF MAJOR HALF AXIS FOR EARTH ELLIPSOID
С
       BQUAD SQUARE OF MINOR HALF AXIS FOR EARTH ELLIPSOID
C ERA, EREQU and ERPOL as recommended by the INTERNATIONAL
C ASTRONOMICAL UNION .
       COMMON/GENER/
                      UMR, ERA, AQUAD, BQUAD
       DATA IENT/0/
       IF (IENT.EQ.0) THEN
           write(*,*)' Initialization call to INITIZE'
           IENT=1
       ENDIF
       ERA=6371.2
       EREQU=6378.16
       ERPOL=6356.775
       AQUAD=EREQU*EREQU
       BQUAD=ERPOL*ERPOL
       UMR=ATAN(1.0)*4./180.
       RETURN
       END
```

С

С

С

C

C

C

C

C

C

С

С C

С

PROGRAM SHIELDFILE_DRIVER This is an auxilliary program to the CREME96 software, which translates a user-supplied shielding distribution into a file with standard format and header information, as required by the CREME96 software. From the user inputs, supplied via interactive dialogue, this program produces an output file. The suggested name of this output file is something.SHD. (ie. the extension should be SHD.) If the file is given some other extension, it will not be accessible by standard CREME96 directory features and (in the WWW version) pull-down menus. IMPLICIT NONE CHARACTER*80 SHIELDFILE, COMMENT CHARACTER*12 MATERIAL INTEGER*4 IUNITS, NBINS, MAXSHIELD PARAMETER (MAXSHIELD=500) REAL*4 XTHICKO (MAXSHIELD), XPROBO (MAXSHIELD)

REAL*4 XTHICK(MAXSHIELD), XPROB(MAXSHIELD) REAL*4 XMEAN, XRMS, TOTAL

INTEGER*4 ERRFLAG

INTEGER*4 VERSION_NUMBER,PROGRAM_CODE

CALL GET_CREME96_VERSION(VERSION_NUMBER) PROGRAM_CODE=7

CALL INISHIELD (MAXSHIELD, COMMENT,

IUNITS, MATERIAL, NBINS, XTHICKO, XPROBO,

SHIELDFILE) &

CALL CHECK_SHIELD_DISTRIBUTION(NBINS, XTHICKO, XPROBO,

XTHICK, XPROB,

XMEAN, XRMS, TOTAL, ERRFLAG)

CALL OUTPUT_SHIELDFILE(SHIELDFILE,

COMMENT, IUNITS, MATERIAL,

NBINS, XTHICK, XPROB,

XMEAN, XRMS, TOTAL, ERRFLAG,

VERSION NUMBER, PROGRAM CODE)

STOP END

&

```
REAL FUNCTION SOLAR_HEAVY_IONS(IZ,EN,IMODEO,ERRFLUX)
C
       Returns the event-integrated interplanetary solar energetic heavy
C
       ion flux (IZ > 2) for element IZ at energy E (in MeV/nuc) for
С
C
       SEP event LSEP.
С
       Inputs:
С
                 = Atomic number (IZ=3-30)
С,
          IZ
                = Energy (MeV/nuc)
С
          IMODE0 = 1: 'worst day' based on measurements of 20 OCT 89 event
С
          IMODE0 = 2: 'worst week' based on measurements of 19-26 OCT 89 events.
C
          IMODE0 = 3: peak flux, based on 5-minute-averaged GOES protons
C
                       on 200CT89
С
С
       Outputs:
C
С
            SOLAR HEAVY_IONS = event integrated flux [in (m2-sr-MeV/nuc)**-1]
C
                    at energy E; NOTE: NOT divided by TIME!
С
            ERRFLUX = its error, based on error propagation of the
C
                    fit parameters. Not yet available.
С
C
       IMPLICIT NONE
       REAL*4 EN, ERRFLUX, FLXDUM
       INTEGER*4 IZ, IMODE0, IMODE, IUSE, IFIT, MELM
       INTEGER*4 NFITS, NMODE, NZ
       PARAMETER (NFITS=2, NMODE=2, NZ=20, MELM=92)
       REAL*4 A1 (NFITS, NMODE), A2 (NFITS, NMODE), A3 (NFITS, NMODE)
       REAL*4 EB1 (NFITS, NMODE), EB2 (NFITS, NMODE)
       REAL*4 BETA (NFITS, NMODE), G(NFITS, NMODE), GAMMA (NFITS, NMODE)
       REAL*4 RELNORM, AVESEP, PEAKFAC, SEP_PEAK_FACTOR
       DIMENSION RELNORM (NZ), AVESEP (MELM)
       DATA A1/ 2.3759E+06, 2.4218E+5, 2.9731E+06, 3.2764E+05 /
       DATA A2/ 4.9518E+08, 1.8991E+08, 1.1307E+09, 3.0372E+08/
       DATA A3/ 0.106702E+10, 0.252948E+10, 0.667628E+09, 0.249719E+09/
       DATA EB1/ 4*0.0 /
       DATA EB2/ 15.94, 24.23, 12.89, 17.22/
       DATA BETA/ 0.5601, 0.2967, 0.4372, 0.2507/
                   5.7000, 5.7000, 5.7000, 5.7000/
        DATA GAMMA/4.14060, 4.52970, 3.76850, 3.7610/
       DATA RELNORM/5*0.0,0.47049,0.12059,1.00000,0.0,
                     0.21312,0.0,0.20624,0.0,0.35935,0.0,
      æ
                     0.09758,0.0,
      &
                     0.00000,0.0,0.04826/
      &
C
      For otherwise undetermined elements Z=6-30, use relative abundances
C
       at 10 MeV/nuc as determined by Croley et al. from the Galileo
C
       data for the 240CT89 event, since these are the best available
C
       observations. AVESEP contains nominal abundances, relative to Fe:
C
C
        DATA AVESEP/5*0.0,
      & 0.4263E+01, 0.1567E+01, 0.1230E+02, 0.5610E-03, 0.1915E+01,
      & 0.2146E+00, 0.3650E+01, 0.2247E+00, 0.2280E+01, 0.2804E-02,
      & 0.1252E+00, 0.2067E-02, 0.2179E-01, 0.4483E-02, 0.9506E-01,
        0.2929E-03, 0.4377E-02, 0.4088E-03, 0.1650E-01, 0.5625E-02,
      & 0.1000E+01, 0.1303E-01, 0.3172E-01, 0.3048E-03, 0.7457E-03,
 C
       For Z>30 elements, include nominal solar abundances as in old CREME.
 C
```

0.4878E-04, 0.1220E-03, 0.7317E-05, 0.7317E-04, 0.9756E-05,

```
The state of the s
```

```
0.9756E-06, 0.4878E-05, 0.0000E+00, 0.2195E-05, 0.4878E-06,
         0.1463E-05, 0.4878E-06, 0.1707E-05, 0.2195E-06, 0.4878E-05,
         0.3415E-06, 0.7317E-05, 0.1463E-05, 0.6585E-05, 0.4878E-06,
         0.4878E-05, 0.4878E-06, 0.1220E-05, 0.1951E-06, 0.9756E-06,
     &
         0.0000E+00, 0.2439E-06, 0.9756E-07, 0.4878E-06, 0.7317E-07,
         0.4878E-06, 0.9756E-07, 0.2439E-06, 0.4878E-07, 0.2195E-06,
         0.4878E-07, 0.1951E-06, 0.2195E-07, 0.2439E-06, 0.4878E-07,
         0.7317E-06, 0.7317E-06, 0.1463E-05, C.2439E-06, 0.2439E-06,
         0.2195E-06, 0.2439E-05, 0.1463E-06, 0.0000E+00, 0.0000E+00,
        0.0000E+00, 0.0000E+00, 0.0000E+00, 0.0000E+30, 0.4878E-07,
         0.0000E+00, 0.2927E-07/
C
       SOLAR HEAVY IONS=0.0
       ERRFLUX=0.0
       IF (EN.LE.O.) RETURN
       IF (IZ.LE.2) RETURN
C
       Set mode flag:
C
       Note: Peak flux (IMODE0=3) is scaled from worst-day (IMODE0=1):
       IMODE=IMODE0
       IF (IMODEO.EQ.3) IMODE=1
       Select baseline spectrum:
       IUSE=8
       IFIT=1
       IF (IZ.GT.20) THEN
                     IUSE=26
                      IFIT=2
       ENDIF
       Nominal modeling:
       IF (EN.LE.EB1(IFIT, IMODE)) THEN
           Note: EB1=0: This segment never activated.
           FLXDUM=A1 (IFIT, IMODE) *EXP(-BETA(IFIT, IMODE) *EN)
       ELSEIF (EB1(IFIT, IMODE).LT.EN .and. EN.LE.EB2(IFIT, IMODE)) THEN
           Forced exponential roll-off at low energies
C
           FLXDUM=EXP(-G(IFIT, IMODE) *EN**0.25)
           FLXDUM=A2(IFIT, IMODE)*FLXDUM*EN**0.25
       ELSEIF (EN.GT.EB2(IFIT, IMODE)) THEN
           Power-law fits at high energies
C
           FLXDUM=A3 (IFIT, IMODE) *EN** (-GAMMA (IFIT, IMODE))
       ENDIF
С
       Special case of broken power-law in worst-week Fe:
       IF (IMODE.EQ.2 .and. IUSE.EQ.26 .and. EN.GT. 127.93) THEN
           FLXDUM=3.168141E+6*EN**(-2.861)
       ENDIF
С
       Scale by relative-abundance factors to get other elements
       IF (IZ.LE.20 .and. RELNORM(IZ).GT.0.) THEN
           SOLAR HEAVY IONS=FLXDUM*RELNORM(IZ)/RELNORM(IUSE)
       ELSE
           SOLAR HEAVY IONS=FLXDUM*AVESEP(IZ)/AVESEP(IUSE)
```

0.4878E-04, 0.7317E-05, 0.2439E-04, 0.4878E-05, 0.1220E-04,

END

```
C
       Finally, convert from /cm2 to /m2:
       SOLAR_HEAVY_IONS=SOLAR_HEAVY_IONS*1.0E+4
С
       For peak-flux model, scale from average:
       IF (IMODEO.EQ.3) THEN
           PEAKFAC=SEP PEAK FACTOR (EN)
           SOLAR_HEAVY_IONS=SOLAR_HEAVY_IONS*PEAKFAC
       ENDIF
       RETURN
       END
       REAL FUNCTION SEP_PEAK_FACTOR (EN)
C
С
       Gets relative scale factor of peak-to-average flux, based on
       GOES proton observations of the 200CT89 SEP event:
С
C
       IMPLICIT NONE
       REAL EN, ENO, ERRFLUX, SOLAR PROTONS, AVEFLUX, PEAKFLUX
       CHARACTER*7 LABEL
       EN0=EN
       IF (ENO.GT.400.) ENO=400.
       LABEL='200CT89'
       AVEFLUX=SOLAR PROTONS (ENO, LABEL, ERRFLUX)
       LABEL='PEAKFLX'
       PEAKFLUX=SOLAR_PROTONS (ENO, LABEL, ERRFLUX)
       SEP PEAK FACTOR=0.0
       IF (AVEFLUX.LE.0.0.or.PEAKFLUX.LE.0.0) RETURN
       SEP_PEAK_FACTOR=PEAKFLUX/AVEFLUX
       RETURN
```

```
C
    C
           Fluences are based on fits to the MEPAD integral proton channels
    C
           on GOES-7 and HEPAD proton channels on GOES-6.
    С
    С
           Inputs:
    С
            E
                  = Energy (MeV/nuc)
    С
             LSEP = SEP event label (CHAR*7, see table below)
    C
    С
             SEP PROTONS = event integrated proton flux [in (m2-sr-MeV/nuc)**-1]
                              at energy E; NOTE: NOT divided by TIME!
    C
    C
                           its error; nominally set to 10%
             SIGFLUX =
    С
    С
    C
           This routine get fluence (differential in energy), starting from
    C
           the fits to the integral rigidity spectrum.
    C
           IMPLICIT NONE
           INTEGER*4 NTERMS, NEVTS, IZ, ISEP
           PARAMETER (NTERMS=4, NEVTS=5)
           REAL*4 E, ERRFLUX, COEF, DAMU, Q, AN, RIGVAL, MAGNETIC RIGIDITY
          REAL*4 XVAL, XDUM, GG, YVAL, FACTOR
           CHARACTER*7 LSEP
           INTEGER*4 K
          DATA DAMU/931.5016/
          DATA IZ/1/,Q/1.0/,AN/1.0/
          DIMENSION COEF (NTERMS, NEVTS)
-
          DATA COEF/
₩
   C 200CT89
         & 0.231924E+02, -0.223621E+02, 0.168443E+02, -0.599497E+01,
   C 190CT89
         & 0.207522E+02, -0.139185E+02, 0.874864E+01, -0.334163E+01,
    C 220CT89
         & 0.215929E+02, -0.155329E+02, 0.790699E+01, -0.258523E+01,
    C 240CT89
         & 0.214273E+02, -0.193048E+02, 0.164647E+02, -0.632318E+01,
    C Peak fluence (/cm2-sr-s) October 1989:
         & 0.135472E+02, -0.232970E+02, 0.185617E+02, -0.674944E+01/
           SOLAR PROTONS=0.0
           ERRFLUX=0.0
           IF (E.LT.1.0 .OR. E.GT.1.0E+5) RETURN
           ISEP=0
          IF (LSEP.EQ.'200CT89') ISEP=1
           IF (LSEP.EQ.'19OCT89') ISEP=2
          IF (LSEP.EQ.'22OCT89') ISEP=3
          IF (LSEP.EQ.'24OCT89') ISEP=4
          IF (LSEP.EQ.'PEAKFLX') ISEP=5
           IF (ISEP.EQ.0) RETURN
             RIGVAL=MAGNETIC RIGIDITY (E,Q,AN)
```

XVAL=COEF(1, ISEP)

DO 500 K=2,NTERMS

XDUM=0.0

REAL FUNCTION SOLAR PROTONS (E, LSEP, ERRFLUX)

Returns the event-integrated interplanetary Solar Energetic Proton

fluence at energy E (in MeV) for SEP event "LSEP" and its estimated

C

C

C

error.

RETURN END

```
GG=FLOAT (K-1)
             XVAL=XVAL+COEF(K, ISEP) *RIGVAL**GG
             XDUM=XDUM+GG*COEF(K, ISEP)*RIGVAL**(GG-1.0)
          CONTINUE
 500
          YVAL=ABS (XDUM) *EXP (XVAL)
       Now need to calculate Jacobian to go from rigidity to kinetic
C
C
       energy:
       FACTOR= (E+DAMU) /SQRT (E*E+2*E*DAMU)
       FACTOR=AN*FACTOR/Q
С
       Additional factor comes from two sources:
С
         1.0E-3 comes from GeV to MeV conversion;
C
         1.0E+4 comes from /cm2 to /m2 conversion.
С
C
       SOLAR PROTONS=10.0*FACTOR*YVAL
C
       For the peak flux mode, the fit parameters give the 5-minute
С
       averaged fluence in (cm2-sr-s)**-1. Need to remove time factor
C
       to put on same basis as other fits:
C
C
       IF (ISEP.EQ.5) SOLAR_PROTONS=SOLAR_PROTONS*300.0
       IF (SOLAR PROTONS.LT.O.) SOLAR_PROTONS=0.0
       ERRFLUX=0.10*SOLAR_PROTONS
```

END

```
SUBROUTINE STABLE (ELOWER, EUPPER, M, IZLO, IZUP, TARGET)
       IMPLICIT NONE
       REAL*4 ELOWER, EUPPER, AA, AMASS, EE, DE, STPOW
       INTEGER*4 M, IZLO, IZUP, NELM, MARR, STAT, CREME96 OPEN
       CHARACTER*12 TARGET
       PARAMETER (MARR=5000, NELM=92)
       REAL*4 SP (NELM, MARR), E (MARR)
       INTEGER*4 J, K, I
       COMMON/MASS/AMASS(109)
С
       Construct list of energies
         DE=(EUPPER/ELOWER) ** (1./(M-1.))
         E(1)=ELOWER
         DO J=2, M-1
           E(J) = E(J-1) *DE
         END DO
         E(M) = EUPPER
          OPEN (UNIT=28, STATUS='NEW', FILE='USER: STABLE.DAT')
C
         stat = creme96 open('stable.dat','user',28,'new')
         WRITE(28,100) ELOWER, EUPPER, M, IZLO, IZUP, TARGET
         WRITE (28, 100)
        DO J=IZLO, IZUP
           AA=AMASS(J)
           DO K=1, M
             EE=E(K)
             SP(J,K)=STPOW(EE,FLOAT(J),AA,TARGET)*AA
           WRITE (28, 200) (SP(J, K), K=1, M)
           Skip line between elements AJT 5/7/96
           WRITE (28, 200)
         END DO
        CLOSE (UNIT=28)
100
      FORMAT (1X, 2 (1PE10.4, 2X), 3 (I5, 2X), A12, 2X, 1PE10.4)
200
      FORMAT((1X,6(1PE10.4,2X)))
      RETURN
```

```
FUNCTION STPOW (E1, Z0, A1, NAME)
         *************
C
            THIS ROUTINE RETURNS THE STOPPING POWER OF NUCLIDE (Z0,A1)
C
           IN MATERIAL 'NAME' AT E1 (MeV/nucleon).
C
           DATA ON THE STOPPING MATERIAL IS CONTAINED IN TARGET.DAT.
C
         ****************
C
        CHARACTER*12 NAME$(150), NAME, LNAME
        REAL IADJ$ (150,28), NA$ (150,28), NASPM$ (150,28)
        INTEGER*4 STAT, CREME96 OPEN
        DIMENSION NAS$ (150), NZ$ (150, 28), DENS$ (150)
        DIMENSION IGAS$(150), ETAD$(150)
        DATA ITARG, LNAME/0, 'QXZ8F3'/
        COMMON /AVG/ AVGZ, AVGZ2, AVGA, AVGI ! MEAN STOPPING MED. PARAMETERS
        * READ IN TARGET DATA
C
        IF (ITARG.EQ.1) GO TO 100
        OPEN(UNIT=10, FILE='CREME96: TARGET.DAT', STATUS='OLD', READONLY, SHARED)
C
        stat = creme96_open('target.dat','cr96tables',10,'old')
 1
        FORMAT(1X, I3)
        FORMAT (1X, A12, 2X, F9.6, 2X, F9.6, 2X, I1, 2X, I2)
        FORMAT (1X, I3, 2X, F8.4, 2X, F5.1, 2X, F9.5)
        READ(10,1) NM$
        DO J1=1, NM$
           \texttt{READ}\,(\texttt{10,2}) \quad \texttt{NAME}\,\$\,(\texttt{J1})\,\,, \texttt{DENS}\,\$\,(\texttt{J1})\,\,, \texttt{ETAD}\,\$\,(\texttt{J1})\,\,, \texttt{IGAS}\,\$\,(\texttt{J1})\,\,, \texttt{NAS}\,\$\,(\texttt{J1})
           DO J2=1, NAS$(J1)
             READ(10,3) NZ$(J1,J2),NA$(J1,J2),IADJ$(J1,J2),NASPM$(J1,J2)
           END DO
         END DO
         ITARG=1
        CLOSE (UNIT=10)
 100
        CONTINUE
        * DETERMINE WHICH TARGET DATA TO USE
         IF (NAME.EQ.LNAME) GO TO 200
        DO J1=1, NM$
           IF (NAME.EQ.NAME$(J1)) K1=J1
        END DO
        LNAME=NAME$(K1)
         IF (NAME.NE.LNAME) THEN
           STPOW=0.
           RETURN
         ENDIF
         * COMPUTE MATERIAL PARAMETERS
С
         RHO=DENS$(K1)
         IGAS=IGAS$(K1)
         ETAD=ETAD$(K1)
         NTOTAL=0
         AVGZ=0.
         AVGZ2=0.
         AVGA=0.
         AVGI=0.
         DO J1=1,NAS$(K1)
           NTOTAL=NTOTAL + NASPM$(K1,J1)
           AVGZ=AVGZ + NASPM$(K1,J1)*FLOAT(NZ$(K1,J1))
           AVGZ2=AVGZ2 + NASPM$(K1,J1)*FLOAT(NZ$(K1,J1))**2
           AVGA=AVGA + NASPM$(K1,J1)*NA$(K1,J1)
           AVGI=AVGI + NASPM$(K1,J1)*ALOG(IADJ$(K1,J1))
         END DO
         AVGZ=AVGZ/FLOAT (NTOTAL)
         AVGZ2=AVGZ2/FLOAT(NTOTAL)
```

```
SUBROUTINE STABLE (ELOWER, EUPPER, M, IZLO, IZUP, TARGET)
           IMPLICIT NONE
           REAL*4 ELOWER, EUPPER, AA, AMASS, EE, DE, STPOW
           INTEGER*4 M, IZLO, IZUP, NELM, MARR, STAT, CREME96 OPEN
           CHARACTER*12 TARGET
           PARAMETER (MARR=5000, NELM=92)
           REAL*4 SP(NELM, MARR), E(MARR)
           INTEGER*4 J, K, I
          COMMON/MASS/AMASS(109)
    C
          Construct list of energies
            DE=(EUPPER/ELOWER) ** (1./(M-1.))
            E(1)=ELOWER
            DO J=2, M-1
               E(J) = E(J-1) *DE
            END DO
            E(M)=EUPPER
    C
             OPEN (UNIT=28, STATUS='NEW', FILE='USER: STABLE.DAT')
            stat = creme96 open('stable.dat','user',28,'new')
            WRITE (28, 100) ELOWER, EUPPER, M, IZLO, IZUP, TARGET
            WRITE (28, 100)
            DO J=IZLO, IZUP
              AA=AMASS(J)
              DO K=1, M
                EE=E(K)
                SP(J,K) = STPOW(EE, FLOAT(J), AA, TARGET) *AA
              WRITE (28, 200) (SP(J, K), K=1, M)
              Skip line between elements AJT 5/7/96
              WRITE (28, 200)
            END DO
            CLOSE (UNIT=28)
100
          FORMAT(1X, 2(1PE10.4, 2X), 3(15, 2X), A12, 2X, 1PE10.4)
   200
          FORMAT ((1X,6(1PE10.4,2X)))
          RETURN
          END
```

```
FUNCTION STPOW(E1, Z0, A1, NAME)
C
        ***************
C
           THIS ROUTINE RETURNS THE STOPPING POWER OF NUCLIDE (ZO,A1)
C
          IN MATERIAL 'NAME' AT El (MeV/nucleon).
C
           DATA ON THE STOPPING MATERIAL IS CONTAINED IN TARGET.DAT.
        ******************
C
        CHARACTER*12 NAME$ (150), NAME, LNAME
        REAL IADJ$ (150,28), NA$ (150,28), NASPM$ (150,28)
        INTEGER*4 STAT, CREME96 OPEN
        DIMENSION NAS$ (150), NZ$ (150, 28), DENS$ (150)
        DIMENSION IGAS$(150), ETAD$(150)
        DATA ITARG, LNAME / 0, 'QXZ8F3' /
        COMMON /AVG/ AVGZ, AVGZ2, AVGA, AVGI ! MEAN STOPPING MED. PARAMETERS
        * READ IN TARGET DATA
C
        IF (ITARG.EO.1) GO TO 100
        OPEN (UNIT=10, FILE='CREME96: TARGET.DAT', STATUS='OLD', READONLY, SHARED)
C
        stat = creme96_open('target.dat','cr96tables',10,'old')
        FORMAT(1X, I3)
 1
        FORMAT (1X, A12, 2X, F9.6, 2X, F9.6, 2X, I1, 2X, I2)
 2
        FORMAT (1X, I3, 2X, F8.4, 2X, F5.1, 2X, F9.5)
 3
        READ(10,1) NM$
        DO J1=1, NM$
          READ(10,2) NAME$(J1), DENS$(J1), ETAD$(J1), IGAS$(J1), NAS$(J1)
          DO J2=1, NAS$ (J1)
            READ(10,3) NZ$(J1,J2),NA$(J1,J2),IADJ$(J1,J2),NASPM$(J1,J2)
          END DO
        END DO
        ITARG=1
        CLOSE (UNIT=10)
 100
        CONTINUE
        * DETERMINE WHICH TARGET DATA TO USE
        IF (NAME.EQ.LNAME) GO TO 200
        DO J1=1, NM$
          IF(NAME.EQ.NAME$(J1)) K1=J1
        END DO
        LNAME=NAMES(K1)
        IF (NAME.NE.LNAME) THEN
          STPOW=0.
          RETURN
        ENDIF
C
        * COMPUTE MATERIAL PARAMETERS
        RHO=DENS$(K1)
        IGAS=IGAS$(K1)
        ETAD=ETAD$ (K1)
        NTOTAL=0
        AVGZ=0.
        AVGZ2=0.
        AVGA=0.
        AVGI=0.
        DO J1=1, NAS$ (K1)
          NTOTAL=NTOTAL + NASPM$(K1,J1)
          AVGZ = AVGZ + NASPM$(K1,J1)*FLOAT(NZ$(K1,J1))
          AVGZ2=AVGZ2 + NASPM$(K1,J1)*FLOAT(NZ$(K1,J1))**2
          AVGA=AVGA + NASPM$(K1,J1)*NA$(K1,J1)
          AVGI=AVGI + NASPM$(K1,J1)*ALOG(IADJ$(K1,J1))
        END DO
        AVGZ=AVGZ/FLOAT (NTOTAL)
        AVGZ2=AVGZ2/FLOAT (NTOTAL)
```

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```
AVGA=AVGA/FLOAT (NTOTAL)
       AVGI=EXP(AVGI/FLOAT(NTOTAL))
       RAT=AVGZ/AVGA
 200
       CONTINUE
       * COMPUTE STOPPING POWERS
C
       STPOW=0.
       IF (E1.LE.O.) RETURN
       EPRIME=0.6443*Z0+13.7144
       XLAMBDA= (E1/EPRIME) **3.2/1.4427
       IF (XLAMBDA.GT.69.) THEN
         WEIGHT=0.
       ELSE
         WEIGHT=EXP (-XLAMBDA)
       ENDIF
       DO J1=1, NAS$ (K1)
         IF (E1.GT.1.) THEN
          TH=RDEDX (E1, Z0, A1, FLOAT (NZ$ (K1, J1)),
          NA$ (K1,J1), IADJ$ (K1,J1),1,1,1,1,RHO, IGAS, ETAD, RAT)
         FLSE
          TH=0.
          WEIGHT=1.
         IF (E1.LT.1000.) THEN
          TL=SPLOW(E1, Z0, A1, FLOAT(NZ$(K1, J1)), NA$(K1, J1))
          TN=SPNUC(E1, Z0, A1, FLOAT(NZ$(K1, J1)), NA$(K1, J1))
        ELSE
          TL=0.
          TN=0.
          WEIGHT=0.
        ENDIF
         TTOTAL=TN + WEIGHT*TL + (1.-WEIGHT)*TH
         STPOW=STPOW+NASPM$(K1,J1)*NA$(K1,J1)*TTOTAL
       STPOW=STPOW/(AVGA*FLOAT(NTOTAL))
       END
     FUNCTION RDEDX(E1, Z0, A1, Z2, A2, IADJ, I0, I1, I2, I3, I4,
    & RHO, IGAS, ETAD, RAT)
C
       AHLEN PROGRAMMED BY SALAMON, MODIFIED BY ADAMS TO
       INCLUDE RELATIVISTIC BLOCH EFFECT.
       CHANGES ARE OUTLINED WITH ASTERISKS.
THIS ROUTINE CALCULATES DE/DX USING THE BETHE EQUATION
C WITH 3 CORRECTION TERMS, THE MOTT, BLOCH, AND LOW VELOCITY Z**3
C TERMS. (SEE S.P. AHLEN, PRA17,1236(1978)). ANY OR ALL OF THESE
C CORRECTIONS CAN BE INCLUDED OR IGNORED, SPECIFIED BY THE
  INDICES I1 (MOTT), I2 (BLOCH), I3 (LOW VELOCITY Z**3). A ZERO INPUT
C FOR A GIVEN PARAMETER ELIMINATES THAT PARTICULAR TERM IN THE
C DE/DX CALCULATION.
C RELATIVISTIC BLOCH CORRECTION (SEE S.P. AHLEN, PRA25,1856(1982))
C IS CONTROLLED BY THE PARAMETER 14 (RELATIVISTIC BLOCH).
C DENSITY EFFECT IS CONTROLLED BY 10 (DENSITY EFFECT).
C
```

```
C THE SHELL CORRECTIONS ARE TAKEN FROM BARKAS AND BERGER,
C PUBLICATION 1133 OF THE NATL ACAD SCI. THE LOW VELOCITY Z**3 CORRECTION
C IS DERIVED FROM A FIGURE IN MCCARTHY AND JACKSON, PHYS REV B6, 4131 (1972).
C THE MOTT, BLOCH CORRECTIONS ARE FROM AHLEN'S PREVIOUSLY
C REFERENCED PAPER.
C
C UNITS OF E1=MEV/AMU
  ZO = ATOMIC NUMBER OF STOPPING NUCLEUS
  Z1 = EFFECTIVE CHARGE OF STOPPING NUCLEUS
C A1 = ATOMIC MASS OF STOPPING NUCLEUS
  Z2 = ATOMIC NUMBER OF ELEMENT IN THE STOPPING MEDIUM.
C
  A2 = ATOMIC MASS OF ELEMENT IN THE STOPPING MEDIUM
C RAT=RATIO OF ELECTRONS/MOLECULE TO NUCLEONS/MOLECULE FOR DENSITY
С
C UNITS OF IADJ=EV. IADJ IS A REAL VARIABLE.
C UNITS OF RETURNED DE/DX=(MEV/AMU)/(G/CM2)
C
C F1=STANDARD DE/DX FRONT FACTOR
C F2=STANDARD BETHE NONRELATIVISTIC TERM WITH SHELL CORRECTIONS
C F3=BLOCH CORRECTION TERM
C F4=LOW VELOCITY Z**3 NONRELATIVISTIC CORRECTION FACTOR
C F5=MOTT CORRECTION TERM (Z**3 TO Z**7)
C F6=STANDARD BETHE RELATIVISTIC TERM
F7=RELATIVISTIC BLOCH CORRECTION TERM
C RHO=DENSITY OF MATERIAL, G/CM**3 (FOR A GAS, GIVE STANDARD DENSITY)
  IGAS=0 IF CONDENSED PHASE, 1 IF GAS.
C ETAD=FOR GAS, DENSITY RELATIVE TO STANDARD (1 ATM, 0 DEG CENT). MUST BE >0.
COMMON/FLOOK/F1, F2, F3, F4, F5, F6, F7, Z1
REAL IADJ
     DIMENSION VA(4), V2FVA(4), Z1ABA(14), COSXA(14)
    DATA Z1ABA, COSXA/0.0,0.05,0.1,0.15,0.20,0.30,0.4,0.5,0.6,
    C 0.8,1.0,1.2,1.5,2.0,1.000,0.9905,0.9631,0.9208,0.8680,
    C 0.7478, 0.6303, 0.5290, 0.4471, 0.3323, 0.2610,
    C 0.2145,0.1696,0.1261/
    DATA VA, V2FVA/1.,2.,3.,4.,0.33,0.30,0.26,0.23/
    PI=3.14159265
    ALPHA=1./137.03604
    G=1.+E1/931.5016
IF IO=O, ELIMINATE DENSITY EFFECT.
      DELT=0.
      IF(I0.EQ.0) GO TO 19
    DELT=DELTA(G, Z2, A2, IADJ, RHO, IGAS, ETAD, RAT)
BSQ=1.-1./G**2
    B=SORT (BSO)
Z1 = Z0
    TEST=B/Z0**(2./3.)
```

```
Z1=Z0*(1.-EXP(-130.*B/Z0**(2./3.)))
          1000 ETA=B*G
          EMASS=0.5110034E+06
                       F1=0.3070722*Z1**2*Z2/(BSO*A2)
                      ETAM2=1./ETA**2
                      CADJ=1.0E-06*IADJ**2*ETAM2*(0.422377+ETAM2*(0.0304043-ETAM2*
                             0.00038106))+1.0E-09*IADJ**3*ETAM2*(3.858019+ETAM2*(-0.1667989
                             +ETAM2*0.00157955))
                      F2=ALOG(2.*EMASS*BSQ/IADJ)-CADJ/Z2
                      F6=2.*ALOG(G)-BSO
                      F3 = 0.0
                      F4=1.0
                      F5=0.0
          IF I2=0, DO NOT CALCULATE BLOCH CORRECTION.
                      IF(I2.EQ.0)GO TO 60
                      Y=Z1*ALPHA/B
                      Y2=Y**2
                     MSUM=INT(5.*Y)+1
                      SUMR=0.
                     DO 90 N=1, MSUM
                     FN=FLOAT(N)
                     FN2=FN**2
         90
                     SUMR = SUMR + (1./(FN2+Y2)-1./FN2)/FN
                     F3 = -Y2 * (1.202 + SUMR)
         C IF I3=0, DO NOT CALCULATE LOW VELOCITY CORRECTION.
         60
                     IF(I3.EO.0)GO TO 50
į.
                     V=ETA/(ALPHA*SQRT(Z2))
9
                      IF(V.GE.4.)GO TO 25
i de la constante de la consta
                     DO 10 I=1,3
IF(V.GE.VA(I+1))GO TO 10
V2FV=V2FVA(I)+(V-VA(I))*(V2FVA(I+1)-V2FVA(I))
12
                     GO TO 30
        10
                     CONTINUE
         25
                     V2FV=0.45/SQRT(V)
        30
                     F4=1.+2.*Z1*V2FV/(V**2*SQRT(Z2))
         C
        C IF I1=0, DO NOT CALCULATE MOTT CORRECTION.
                     IF(I1.EQ.0)GO TO 70
                     Z1A=Z1*ALPHA
                     Z1AB=ABS (Z1A/B)
                     COSX=0.
                     DO 40 I=1,13
                     IF(Z1AB.GE.Z1ABA(I+1))GO TO 40
                     {\tt COSX=COSXA(I)+(Z1AB-Z1ABA(I))*(COSXA(I+1)-COSXA(I))/}
                   C (Z1ABA(I+1)-Z1ABA(I))
        40
                     CONTINUE
                     F5=0.5*Z1A*(B*(1.725+0.52*PI*COSX)+Z1A*(3.246-0.451*BSQ
                           +Z1A* (1.522*B+0.987/B+Z1A* (4.569-0.494*BSQ-2.696/BSQ
                              +Z1A*(1.254*B+0.222/B-1.170/BSQ/B)))))
                     IF(Z1AB.LE.100.*ALPHA)GO TO 70
        ERR=Z1AB**9/6.
                     IF (ERR .GT. .01) ERR=.01
                     IF(ERR.LT.ABS(F5/(F2*F4+F3+F5+F6-DELT/2.)))GO TO 70
```

IF (TEST .GT. .1) GO TO 1000

```
C C C C 10 C 10 C 21 11 20 21 100 C
```

```
IF I4=0, DO NOT CALCULATE RELATIVISTIC BLOCH CORRECTION.
70
        IF(I4.EQ.0)GOTO 80
        F7=CR(Z1,G,B)
      RDEDX=F1*(F2*F4+F3+F5+F6+F7-DELT/2.)/A1
80
      END
      FUNCTION DELTA (G, Z2, A2, FIADJ, RHO, IGAS, ETA, RAT)
   THIS FUNCTION IS USED BY RDEDX.
  THIS CORRECTION FOR THE DENSITY EFFECT IS BASED ON STERNHEIMER AND
C
  PEIERLS, PHYS REV B3, 3681 (1971).
   SET IGAS = 0 OR 1, ETA > 0 REQUIRED.
   RHO IS DENSITY IN G/CM**3. FOR A GAS, GIVE RHO AT T=0 DEGREES
   CENTIGRADE, 1 ATM PRESSURE, AND THE FACTOR ETA WHICH GIVES THE
  ACTUAL GAS DENSITY UPON MULTIPLICATION BY RHO.
      IF (G.GE.1.8) GO TO 10
      DELTA=0.
      RETURN
        *****************
      PLASMA=28.8*SQRT(RHO*RAT) ! RAT REPLACES Z2/A2 FOR MOLECULE
10
      CBAR=2.*ALOG(FIADJ/PLASMA)+1.0
      B = SQRT(1.-1./G**2)
      Y=2.*ALOG(B*G)+IGAS*ALOG(ETA)
      IF(IGAS.EQ.1)GO TO 100
      IF(FIADJ.GE.100.)GO TO 20
      Y1 = 9.212
      IF(CBAR.GE.3.681)GO TO 11
     Y0 = 0.9212
      GO TO 200
     Y0=1.502*CBAR-4.606
     GO TO 200
     Y1=13.82
      IF(CBAR.GE.5.215)GO TO 21
     Y0=0.9212
     GO TO 200
     Y0=1.502*CBAR-6.909
     GO TO 200
100
     IF(CBAR.GE.12.25)GO TO 110
     Y1=18.42
     IF(CBAR.LT.12.25)Y0=9.212
     IF (CBAR.LT.11.5) Y0=8.751
     IF (CBAR.LT.11.0) Y0=8.291
     IF(CBAR.LT.10.5)Y0=7.830
     IF (CBAR.LT.10.0) Y0=7.370
     GO TO 200
110
     Y1 = 23.03
     IF(CBAR.GE.13.804)GO TO 120
     Y0 = 9.212
     GO TO 200
120
     Y0=1.502*CBAR-11.52
200
     A = (CBAR - Y0) / (Y1 - Y0) **3
```

```
IF(Y.GT.Y0)GO TO 210
DELTA=0.
RETURN

210 IF(Y.GE.Y1)GO TO 220
DELTA=Y-CBAR+A*(Y1-Y)**3
RETURN

220 DELTA=Y-CBAR
RETURN
END
```

FUNCTION SIGMA (NU)

```
C
        **********************
C
        ** THIS FUNCTION COMPUTES THE PHASE OF THE GAMMA FUNCTION OF
C
        ** 1 + i*NU. IT IS CALLED BY THE FUNCTION CR WHICH COMPUTES
С
        ** THE RELATIVISTIC BLOCH CORRECTION.
C
        ***********************
        REAL*4 NU
        DIMENSION A(7), H(7)
        DATA(A(I), I=1,5)/.26356,1.4134,3.59642,7.0858,12.6408/
        DATA(H(I), I=1,5)/.521756,.398667,.075942,.003612,23E-6/
        SUM=0.
        DO 1 I=1,5
        SUM=SUM+H(I)*SIN(NU*ALOG(A(I)))
 1
        CONTINUE
        DEM=0.
        DO 2 I=1,5
       DEM=DEM+H(I) *COS(NU*ALOG(A(I)))
 2
        CONTINUE
        SIGMA=ATAN2 (SUM, DEM)
       RETURN
       END
       FUNCTION CR (Z, GAMMA, BETA)
        ********************************
C
C
        ** THIS FUNCTION COMPUTES THE RELATIVISTIC BLOCH
        ** CORRECTION TO THE STOPPING POWER ACCORDING TO THE WORK
C
C
        ** OF S.P. AHLEN, PRA25, 1856 (1982).
C
       REAL*4 LAMBDA, NU, IL1, LNROW, LNROWT
       DATA EULER/.577215665/,PI/3.14159265/
       DATA AMU/931.5016/, ALPHA/.0072973504/
       DATA LAMBDA/1.0/, THETA/0.1/
C
    DEFINITIONS
       NU=Z*ALPHA/BETA
       TNU=2.*NU
       DEM=1./(1.+TNU*TNU)
       TNLT=TNU*ALOG(0.5*THETA)
       TNT=TNU*THETA
       TODEM=THETA*DEM
C
    COMPUTE REAL AND IMAGINARY PARTS OF L1
       RL1=TODEM* (SIN (TNLT) - TNU*COS (TNLT))
       IL1=-TODEM* (COS (TNLT) +TNU*SIN (TNLT))
       AOBGL=ALPHA/(BETA*GAMMA*LAMBDA)
       TNLOG=TNU*ALOG (AOBGL)
```

```
ARG=TNLOG+2.*SIGMA(NU)
          Bug here discovered by Bonnie Colborn 7-7-95
   C
   C
          RL1=RL1+COEF* (TNU*COS (ARG) -SIN (ARG) )
          IL1=IL1+COEF* (TNU*SIN(ARG)+COS(ARG))
   C
          RL1=RL1+COEFF* (TNU*COS (ARG) -SIN (ARG))
          IL1=IL1+COEFF* (TNU*SIN(ARG)+COS(ARG))
        COMPUTE REAL PART OF L2
   C
          FNLOG=2.*TNU*DEM-TNLT
          FNOLOG= (TNU*TNU-1) *DEM+ALOG (0.5*THETA)
          RL2=TODEM* (FNLOG*COS (TNLT) +FNOLOG*SIN (TNLT))
          LNROW=ALOG(2./AOBGL)+EULER-1+(1-TNU*TNU)*DEM
          LNROWT=ALOG(2./AOBGL)+EULER-1+2*DEM
          RL2=RL2+COEFF* (LNROW*SIN(ARG)-TNU*LNROWT*COS(ARG))
   C
        COMPUTE CR
          PINU=PI*NU
          CR=0.5*PINU*BETA*BETA* (PINU*EXP(PINU) /SINH(PINU))
          CR=CR*(2.*TNU*ALOG(2.)*RL1+(PINU-1.)*IL1+TNU*RL2)
          RETURN
          END
          FUNCTION HYDRGN (EN, Z2, A2)
          **************
   С
   C
            STOPPING POWER OF SLOW PROTONS (1 KeV to 1 MeV)
          ******************
   C
          INTEGER*4 STAT, CREME96_OPEN
          DIMENSION A (92, 12)
          DATA MARKER/0/
          IZ2 = INT(Z2 + 0.2)
          IF (Z2.GT.92.) IZ2=92
          ******************
į.
   C
            ON FIRST CALL TO FUNCTION (MARKER=0) READ IN DATA FROM
æ
          * PROTON.DAT.
   C
   C
          ************
          IF (MARKER.EQ.0) THEN
            OPEN (UNIT=50, READONLY, STATUS='OLD', FILE='CREME96: PROTON.DAT', SHARED)
            stat = creme96 open('proton.dat','cr96tables',50,'old')
            DO I=1,92
              READ(50,20) (A(I,J),J=1,11)
   20
              FORMAT (11 (1X, E10.4))
            END DO
            CLOSE (UNIT=50)
            MARKER=1
          ENDIF
          E=EN*1.007825*1000. ! CHANGE FROM MEV/NUCLEON TO KEV
          **********************
   Ċ
   C
            COMPUTE STOPPING POWER IN (MEV/NUCLEON)/(G/CM**2)
   C
          *******************
          IF (E.LE.O.) THEN
            HYDRGN=0.
            RETURN
          ELSE IF (E.LT.1000.) THEN
            SL=A(IZ2,1)*E**.45
            SH = (A(IZ2,2)/E)*ALOG(1.+A(IZ2,3)/E+A(IZ2,4)*E)
            S=SL*SH/(SL+SH)
          ELSE
            G=1.+EN/931.5016
            BSQR = (1.-1./(G*G))
            COEFF=A(IZ2,5)/(BSQR)
```

COEFF=AOBGL*DEM

```
35
ĦŲ.
```

C

C C

```
SHELL=ALOG(A(IZ2,6)*BSQR/(1.-BSQR))-BSQR
         ALOGE=ALOG(E)
         SHELL=SHELL-(A(IZ2,7)+A(IZ2,8)*(ALOGE))
         SHELL=SHELL-(A(IZ2,9)*((ALOGE)**2.)+A(IZ2,10)*((ALOGE)**3.))
         SHELL=SHELL-(A(IZ2,11)*((ALOGE)**4.))
         S=COEFF*SHELL
       ENDIF
       HYDRGN=S*1.E-21/(A2*1.007825*1.659828E-24)
       IF (Z2.GT.92.) HYDRGN=HYDRGN*(Z2**2/92.**2)
       RETURN
       END
       FUNCTION HELIUM (EN, Z2, A2)
C
       **********************
C
          STOPPING POWER OF SLOW ALPHAS (1 KEV TO 1 MEV)
       *******************
C
       INTEGER*4 STAT, CREME96 OPEN
       DIMENSION A (92,9)
       DATA MARKER/0/
       IZ2=INT(Z2+0.2)
       IF (Z2.GT.92.) IZ2=92
       ***********
C
C
         ON FIRST CALL TO FUNCTION (MARKER=0) READ IN DATA FROM
С
       * HELIUM.DAT.
C
       ***********
       IF (MARKER.EO.O) THEN
        OPEN (UNIT=40, READONLY, STATUS='OLD', FILE='CREME96: HELIUM.DAT', SHARED)
        stat = creme96 open('helium.dat','cr96tables',40,'old')
        DO I=1,92
C
          READ (40,20) (A(I,J),J=1,9)
C20
          FORMAT (9(1X,E10.4))
          READ (40,20) (A(I,J),J=1,9)
          FORMAT (1X, 9 (E9.4))
        END DO
        CLOSE (UNIT=40)
        MARKER=1
      ENDIF
      E=EN*4.0026*1000. ! CHANGE FROM MEV/NUCLEON TO KEV
      *************
      * COMPUTE STOPPING POWER IN (MEV/NUCLEON)/(G/CM**2)
      IF (E.LE.O.) THEN
        HELIUM=0.
        RETURN
  CHANGE IN UPPER ENERGY LEVEL, R.A. WITT 17 MARCH 1994
      ELSE IF (E.LT.10000.) THEN
      ELSE IF (E.LT.100000.) THEN
        SL=A(IZ2,1)*E**A(IZ2,2)
        SH=(A(IZ2,3)/(E/1000.))*ALOG(1.+A(IZ2,4)/(E/1000.)+A(IZ2,5)
        *(E/1000.))
        S = (SL*SH/(SL+SH))
      ELSE
        EE=ALOG(1/(E/1000.))
        S=EXP(A(IZ2,6)+A(IZ2,7)*EE+A(IZ2,8)*EE*EE+A(IZ2,9)*EE*EE*EE)
      ENDIF
```

```
END
       FUNCTION SPLOW(E, Z0, A1, Z2, A2)
       *******************
C
          STOPPING POWER OF SLOW NUCLEI
C
       C
       IZ1=INT(Z0+0.2)
       IF (IZ1.EQ.1) THEN
         SPLOW=HYDRGN (E, Z2, A2) *1.007825/A1
       ELSE IF (IZ1.EQ.2) THEN
         SPLOW=HELIUM (E, Z2, A2) *4.0026/A1
       ELSE
         C=2.99792458E10
         V=C*(1.-(1./(E*1.6022E-6/C/1.673E-24/C+1.))**2.)**0.5
         V1=0.886*(V/2.188E8)*Z0**(-2./3.)
         V2=V1+0.0378*SIN(V1*3.14159265/2.)
         SCALE=(1.-(1.034-0.1777*EXP(-0.08114*Z0))*EXP(-V2))**2.
         SPLOW=SCALE*Z0**2*HYDRGN(E, Z2, A2)*1.007825/A1
       ENDIF
       RETURN
       END
       FUNCTION SPNUC (E, Z0, A1, Z2, A2)
С
       **************
C
        LNS STOPPING POWER AS PROGRAMMED BY JR LETAW
       **************
C
       DATA ZE, AO, EC/4.803242E-10, 5.2917706E-9, 1.6021892E-6/
       DATA AV, PI, XL/6.022045E23, 3.1415927, 1.309/
       DATA T1/4.005473E-11/ ! MIN. E = 25 eV
       SPNUC=0.
       IF ((E*A1).LT.2.5E-5) RETURN
       Z = (Z0**(2./3.) + Z2**(2./3.)) **1.5
       A=0.8853*A0/Z**(1./3.)
       EL=Z0*Z2*(ZE**2/A)*(A1+A2)/A2
       G=4.*A1*A2/(A1+A2)**2
       ECONV=E*A1*EC
       P1=(2.*XL)**(1./3.)*(ECONV*T1/EL**2/G)**(2./9.)
       P2 = (2.*XL)**(1./3.)*(ECONV/EL)**(4./9.)
       FACT=1.125*PI*A**2*(EL**2*G/ECONV)*(AV/A2)/A1/EC
       P1S=SQRT(1.+P1**2)
       P2S=SQRT(1.+P2**2)
       SPNUC=ALOG(P2+P2S)-P2/P2S
       SPNUC=FACT* (SPNUC-(ALOG(P1+P1S)-P1/P1S))
       RETURN
       END
```

HELIUM=S*1.E-21/(A2*4.0026*1.659828E-24)
IF (Z2.GT.92.) HELIUM=HELIUM*(Z2**2/92.**2)

RETURN

```
C**********************************
C Special version of UPROPI, for doing transport through thin shield
C without utilizing external files of dE/dx and range-energy, as generally
C done in the UPROP routines.
C Important variables
С
CE
           Energy at each grid point after shielding
C S
           Stopping power at each grid point after shielding
C R
           Range at each grid point after shielding
C EP
           Energy at each grid point prior to shielding
C SP
           Stopping power at each grid point prior to shielding
C RP
          Range at each energy EP
C
      IMPLICIT NONE
      INTEGER*4 MARR, NELM
      PARAMETER (MARR=5000, NELM=92)
      REAL*4 FLUX (NELM, MARR), E (MARR), FLUX2 (MARR)
      REAL*4 R (MARR), S (MARR), EP (MARR), RP (MARR), SP (MARR)
      CHARACTER*12 TARGET
      INTEGER*4 M, IZLO, IZUP, J, K, L, KK, LMAX
      REAL*4 ELOWER, EUPPER, PATH, REL, FUL, DE, XK, AMASS, Z, A
      REAL*4 STPOW
      COMMON/MASS/AMASS(109)
      DATA LMAX/2/
      Compute vector of energies
      REL=1./ELOWER
      FUL=1./LOG(EUPPER/ELOWER)
      DE=(EUPPER/ELOWER) ** (1./FLOAT(M-1))
      E(1) = ELOWER
      DO J=2, M-1
        E(J) = E(J-1) *DE
      END DO
      E(M)=EUPPER
C
      Compute range-energy relations and stopping powers:
      DO J=IZLO, IZUP
        Z=FLOAT(J)
        A=AMASS(J)
        CALL RANGE (E, M, Z, A, TARGET, R)
        DO K=1,M
          S(K) = STPOW(E(K), Z, A, TARGET)
        END DO
        DO K=1, M
          DO KK=K, M
             IF (R(KK).GE.R(K)+PATH) GOTO 300
          END DO
          KK=M
300
          EP(K) = E(KK) - (R(KK) - R(K) - PATH) *S(KK)
          R(K) = R(K) + PATH
        END DO
```

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```

```
C
         Iterate LMAX times to improve estimate of EP
        DO L=1, LMAX
           CALL RANGE (EP, M, Z, A, TARGET, RP)
           DO K=1, M
             SP(K) = STPOW(EP(K), Z, A, TARGET)
             EP(K) = EP(K) - (RP(K) - R(K)) *SP(K)
           END DO
        END DO
C
С
        Now get flux values at these corresponding external energies:
        DO K=1, M
          XK=1.+(M-1.)*LOG(EP(K)*REL)*FUL
          KK=INT(XK)
           IF (XK.GE.M) THEN
             FLUX2(K) = ((EP(K) - E(M-1)) * FLUX(J,M) +
     &
                       (E(M)-EP(K))*FLUX(J,M-1))/(E(M)-E(M-1))
           ELSE
             FLUX2(K) = ((EP(K) - E(KK)) * FLUX(J, KK+1) +
                       (E(KK+1)-EP(K))*FLUX(J,KK))/(E(KK+1)-E(KK))
          ENDIF
          FLUX2(K) = FLUX2(K) *SP(K) /S(K)
          IF (FLUX2(K).LT.1.E-20) FLUX2(K)=0.
        END DO
        DO K=1, M
          FLUX(J,K) = FLUX2(K)
        END DO
      END DO
      RETURN
      END
```

PROGRAM TRANSPORT DRIVER

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LJ

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Now write transported flux to output file:

CALL OUTPUT_TRANSPORTED_FLUX(IZLO,IZUP,ELOWER,EUPPER,

IPATH, UPATH, TARGET,

SHIELDFILE, INFILE,

VERSION_NUMBER, PROGRAM_CODE,

M, OUTPUT_FLUX, OUTFILE)

STOP END

С

С

```
subroutine trapped_protons(B, L, yearp, jmod, energy, flux, ne)
                *******************
C
       Inputs:
С
           B from Blccoords
C
           L from Blccoords
C
           yearp from Blccoords
           jmod, = 1 for solar min model, = 2 for solar max model
С
C
           energy, an array of values in MeV
C
           ne, the number of energy values in the energy array
C
       Outputs:
           flux, ne values of integral flux greater than the corresponding
C
C
                     Mev value in the energy array
implicit none
      save
      real*4 a8max, a8min, B, energy, flux, L, yearp,f1, fi1
      integer*4 ie, ifirst, itpfile, jmod, jmodold, 18min, 18max
      integer*4 ne, map
      dimension energy(1), flux(1)
      real*8 gmagmo
      common /gmagmo/ gmagmo
                                        for esa traraln
                                   1
      common /energy/ f1(30,45),fi1(30,8)
      common /sumry/ map(777)
      common/ap8min/a8min(8),18min(16583)
      common/ap8max/a8max(8),18max(16583)
      integer CREME96 OPEN, stat
      data ifirst /0/
      data itpfile /8/
      if (ifirst .eq. 0 ) then
        ifirst = 1
           jmodold = jmod
                             !jmod =1 for min =2 for max
           if(jmod .lt. 1 .or. jmod .gt. 2) stop 'tp model1'
           stat = creme96_open('ap8maxmi.inp','cr96tables',itpfile,
                               'old')
C
        read in the proton model data to be used
        call modint(itpfile,a8min,18min)
        read(itpfile, 16)
        call modint(itpfile,a8max,18max)
16
        format(19a4)
        close(unit=itpfile)
     endif
 500 continue
     if (jmod .ne. jmodold) then
         type *, ' Model number input has changed '
          stop 'tp model2'
     endif
     if ((b.eq. 0.) .or. (L.eq. 0.)) return
                                             !not in range of values
     if ( L .gt.11.) return ! not in range of values
     if (jmod .eq. 1) then
         call traral(a8min, 18min, L, B, energy, flux, ne)
     elseif (jmod .eq. 2) then
         call traral(a8max, 18max, L, B, energy, flux, ne)
```

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```
else
         type *, ' Illegal model number input to trapped_protons'
         stop 'tp model3'
     endif
     do ie = 1, ne
        flux(ie) =10.**flux(ie)
        if (flux(ie) .lt. 1.001) flux(ie) = 0.
     enddo
     return
     end
C*********************************
     subroutine modint(junit, descr, list)
implicit none
     save
     dimension descr(8), list(16583)
     real *4 descr, dumd
     integer*4 junit, list, length, ic, lnt, i, jc
     integer*4 k, k1, k2, lb, lp, lpp
     equivalence (length, dumd)
     read (junit, 1000, end=30) (descr(i), i=1,7), length, lb, ic
     descr(8) = dumd
     type 1002, (descr(i), i = 1,7), length, lb, ic
     lnt = length+1
     lp = lnt/7
     lpp = lp*7
     if (lpp.ne.lnt) lp = lp+1
     lp = lp+1
     k1 = 1
     do jc = 2, lp
       k2 = k1+6
       read(junit, 1001, end=30)(list(k), k=k1, k2), lb, ic
       k1 = k2+1
     enddo
     return
  30 type*, ' *** read eof on ',junit,' ***'
     stop 'modint'
1000 format (2a4, 2x, 5f10.3, i10, a4, i4)
1001 format (7i10, a4, i4)
1002 format(1x, 2a4, 2x, 5f10.3, i10, 2x, a4, i4)
subroutine traral(descr, map, fl, babs, e, f, n)
B / BO CASE
C
                   JOEL STEIN
                                9-15-71
                                          X2133
  TRARAL DOES ENERGY VALUE SEARCH FOR FLUX CALCULATION WHEN GIVEN A
C
    B AND L POINT.
* Modified version based on Kluge and Lenhart ESOC Int. Note 78 (1971)
             APRIL 1988 E.J. DALY ESA/ESTEC/WMA
  Modified
                      can pick up geomagnetic dipole moment GMAGMO
                       from common block and use it in place of
                      McIlwain's value of .311653
                      GMAGMO is passed from the geomagnetic
                      coordinate program.
                      Default for models should be McIlwain's value
```

```
Modified
                 JULY 1993 H.D.R. EVANS
                                            ESA/ESTEC/WMA
                             Kluge and Lenhart interpolation method in
                             (B/B0,L) space changed to a linear polygon
                             interpolation method in (Phi,L) space.
   C MAP(1) is the first word of list
   C S0,S1,S2 are logical variables which indicate whether the flux for a
   C particular E,B,L point has already been found in a previous call
   C to TRARAP.
         implicit none
         save
         logical s0,s1,s2
         real*4 bob0, babs, descr, e, e0, e1, e2, f, f1, f0, f1, f2
         real*4 trarap
         integer*4 ie, i0, i1, i2, i3, l3, map, nb, n, nl
         dimension e(1), f(1), descr(8), map(1)
         real*8 gmagmo
         common / gmagmo / gmagmo
nl=amin1(32766.,abs(fl*descr(5)))
   C MAX B/B0 ALLOWED HERE IS 1000 (PROTECT AGAINST INTEGER OVERFLOW)
         if (gmagmo .le. 0.) gmagmo = 0.311653
C
        BOBO = AMIN1 ( (BABS*(FL*FL*FL) / GMAGMO ), 1000.)
į.
        bob0 = (babs*(fl*fl*fl) / gmagmo)
         if (bob0 .gt. 1000.) bob0 = 1000.
i.i.
TI C
        HANDLE CASE WHERE B/B0 IS LESS THAN 1.0 (DISREGARDING REPRESENTATIONAL
LJ c
        errors)
i.i.
         if (bob0 .lt. 0.95) then
             do ie = 1, n
f(ie) = -99.0 ! CHECK TO SEE IF -99 IS USED IN CALLING ROUTINE??????
             enddo
             return
        endif
   C
        FORCE ANY POSSIBLE REPRESENTATIONAL ERRORS TO 1.0
        BoB0 = AMAX1(BoB0, 1.0)
        nb = abs((bob0-1) * descr(6))
   C NL IS THE MINIMUM OF THE L VALUE OR 15.999, SCALED TO AN INTEGER BY
        THE L SCALING FACTOR
   C
      NB IS THE DIFFERENCE BETWEEN THE INPUT B VALUE AND B EQUATORIAL,
        SCALED TO AN INTEGER BY THE B SCALING FACTOR.
        i1 = 0
        i2 = map(1)
        i3 = i2 + map(i2 + 1)
        13 = map(i3+1)
        e1 = map(i1+2)/descr(4)
        e2 = map(i2+2)/descr(4)
        s1 = .true.
```

39

s2 = .true.

12 IS THE NUMBER OF ELEMENTS IN THE FLUX MAP FOR THE FIRST ENERGY. I3 IS THE INDEX OF THE LAST ELEMENT OF THE SECOND ENERGY MAP. C L3 IS THE LENGTH OF THE MAP FOR THE THIRD ENERGY. C E1 IS THE ENERGY OF THE FIRST ENERGY MAP (UNSCALED) C E2 IS THE ENERGY OF THE SECOND ENERGY MAP (UNSCALED) C S1 AND S2 ARE TRUE TO INDICATE THAT NO FLUXES HAVE YET BEEN FOUND. do 3 ie = 1,nTHE DO STATEMENT LOOPS THROUGH THE ENERGIES FOR WHICH FLUXES ARE C DESIRED AT THE GIVEN B, L POINT (BABS, FL). 1 if (e(ie).le.e2.or.13.eq.0)goto2 THE IF STATEMENT CHECKS TO SEE IF THE INPUT ENERGY IS LESS THAN OR E THE ENERGY OF THE SECOND MAP, OR IF THE LENGTH OF THE THIRD MAP IS C (I.E. THERE ARE NO HIGHER ENERGIES IN THE TABLE). IF TRUE, USE TH FOR THOSE TWO ENERGY MAPS TO FIND THE DESIRED FLUX AT THE DESIRED C ENERGY. IF FALSE, THE ZEROTH ENERGY MAP IS DEFINED TO BE THE FIRS C C ENERGY MAP, THE FIRST BECOMES THE SECOND, AND THE SECOND BECOMES THE THIRD. E0, E1, E2 ARE THE ENERGIES FOR THE ZEROTH, FIRST, AND SEC С ENERGY MAPS. F0,F1,F2 ARE THE FLUXES FOR THE ZEROTH, FIRST, AND SECOND ENERGY MAPS AT THE B,L POINT. i0 = i1i1 = i2i2 = i3i3 = i3+1313 = map(i3+1)e0 = e1e1 = e2e2 = map(i2+2)/descr(4)s0 = s1s1 = s2s2 = .true.f0 = f1f1 = f2goto1 if (s1) f1 = trarap(descr, map(i1+1), f1, bob0) if (s2)f2 = trarap(descr, map(i2+1), fl, bob0) C THESE TWO LOGICAL IFS CALL TRARAP FOR THE FLUX FROM THE FIRST AND SECOND ENERGY MAPS AT THE B,L POINT IF THEY HAVE NOT ALREADY BEEN s1 = .false.S2 = .FALSE.C S1 AND S2 ARE FALSE SINCE F1 AND F2 ARE NOW FOUND. f(ie) = f1+(f2-f1)*(e(ie)-e1)/(e2-e1)INTERPOLATE FOR THE FLUX F(IE) USING THE FLUXES AND ENERGIES FOR MAP C C ONE AND TWO. THE FOLLOWING COMMENTS APPLY TO THE REMAINING PROGRAM STATEMENTS. C C IF THE FLUX F2 FOR THE SECOND ENERGY MAP IS GREATER THAN ZERO, OR TH ZEROTH ENERGY MAP HAS NOT BEEN DEFINED, THE FINAL FLUX IS THE MAXI C OF THE INTEROOLATED FLUX OR ZERO. IF THE FLUX FOR THE SECOND ENER C MAP IS EQUAL TO ZERO, AND THE ZEROTH ENERGY MAP HAS BEEN DEFINED. C THEN INTERPOLATE FOR THE FLUX USING THE ZEROTH AND FIRST ENERGY MA CHOOSE THE MINIMUM OF THE TWO INTERPOLATIONS, AND THEN THE MAXIMUM C CHOICE AND ZERO FOR THE FINAL FLUX VALUE.

```
if(f2.qt.0.)goto3
     if(i1.eq.0)goto3
     if(s0)f0 = trarap(descr, map(i0+1), fl, bob0)
     s0 = .false.
     f(ie) = amin1(f(ie), f0+(f1-f0)*(e(ie)-e0)/(e1-e0))
     f(ie) = amax1(f(ie), 0.)
     return
     end
REAL FUNCTION TRARAP ( HEADER, MAP, L, BoB0)
This function converts the B argument to a hybrid
С
     PURPOSE:
                value, defined by Phi = ASIN( ( B - B0) / (Bmax- B0))
C
                and interpolates the MAP in (PHI-L) space.
C
C
                The conversion to the Phi-L space requires the
     METHOD:
С
                maximum B value. This is obtained by interpolating
С
                between the maximum B values for the L strings
С
                that subtend the L value passed to the routine.
C
C
                The Flux at the Phi point along the two L strings
C
                is determined by locating the three model points
С
                that form a polygon (triangle) that contains the
С
                (L,Phi) point. The interpolation is then performed
C
                by a linear interpolation using the slope of the
C
                plane in (L, Phi, lnF) space.
C
C
                                  H.D.R. Evans ESA/ESTEC/WMA
               CREATED July 1993
     HISTORY:
implicit none
     save
     real bob0, 1, b, phi
                map(*)
     integer*4
                header(*)
     real
              model(3,100,2)
     real
               ls(100), bs(100), fluxes(100)
     real
               bm(2), bo
     real
                verts(3,3)
     real
                reqpt(3)
     real
               xphi, b0, xinter, bmax, interp
     real
     integer*4
                estrng, lstrng
                  lpos(2), llen, blen(2), li
     integer*4
                 i, j
     integer*4
                 s(2), n(2), string, other, tmp
     integer*4
                found, endstr
     logical
     logical
                inpoly
     if (bob0 .lt. 1.0) then
        trarap = 0.0
        return
      endif
     b = bob0 * 0.311653 / (1**3)
                                     !????ASK TONY: SHOULD WE USE GMAGMO?
      call lvals ( map, header, ls, llen)
      FIND THE TWO L STRINGS THAT SUBTEND THE L VALUE WE ARE GIVEN...
 C
      do li = 1, llen-2
        if ( l .le. ls(li+1) ) go to 6
      enddo
      continue
```

```
GET THE MAXIMUM B VALUES FOR THE TWO L STRINGS. ALLOWS US TO
С
      LINEARLY INTERPOLATE TO FIND THE MAXIMUM B VALUE FOR THE GIVEN
С
      L VALUE.
C
      do i = 1, 2
         lpos(i) = lstrng( map, header, ls(li+i-1), llen) + 1
         call bvals ( map( lpos(i)), header, bs, fluxes,
         bm(i) = bs(blen(i))
         bo = b0 ( ls(li+i-1))
         if (bo .gt.0.) then
            do j=1, blen(i)
               model(1,j,i) = ls(li+i-1)
               model(2,j,i) = xphi(bs(j)/bo, bm(i) / bo)
               model(3,j,i) = fluxes(j)
            enddo
         else
            blen(i) = 1
            model(1,1,i) = ls(li+i-1)
            model(2,1,i) = -1.0
            model(3,1,i) = 0.0
         endif
      enddo
      bo = b0(1)
      phi = xphi( b / bo, xinter( l, ls(li),
                                     ls(li+1), bm(2)) / bo)
     &
      CHECK FOR AN INVALID PHI VALUE, E.G. IF B > BMAX
      if (phi .lt. 0 ) then
         trarap = 0.0
         return
      endif
      IF LENGTH OF BOTH STRINGS IS 1, THEN LINEAR INTERPOLATION BETWEEN
C
      POINTS AND RETURN
      if (blen(1) .eq. 1 .and. blen(2) .eq. 1) then
                         ( l - model(1,1,1)) *
         trarap =
                          ( model(3,1,2) - model(3,1,1) ) /
      æ
                          ( model(1,1,2) - model(1,1,1) )
         if (trarap .gt.0) trarap = ( trarap)
         return
      endif
      NOW FIND H VALUES IN BOTH L STRINGS THAT SUBTEND REQUIRED POINT.
C
      reqpt(1) = 1
      reqpt(2) = phi
       s(1) = 1
       s(2) = 1
      n(1) = min(s(1)+1, blen(1))
      n(2) = min(s(2)+1, blen(2))
  10
      continue
       endstr = (s(1) .eq. blen(1)) .and. (s(2) .eq. blen(2))
       if (.not. endstr) then
          if (s(1) .eq. blen(1)) then !string 1 is empty, have to use string 2 now.
             string = 2
```

C

```
other = 1
       else if (s(2).eq.blen(2)) then! string 2 is empty, have to use string 1.
          string=1
          other =2
       else
          if ( model(2, n(1), 1) .lt. model(2, n(2), 2) ) then
             string = 1
             other = 2
          else
             string = 2
             other = 1
          endif
       endif
       found = inpoly( model(1, s(string), string),
                               model(1, s(other), other),
   &
                               model(1, n(string), string), reqpt )
       if ( .not. found ) then
          s(string) = n(string)
          n( string) = min( n(string) +1, blen( string) )
       endif
    endif
    if (.not. (found.or.endstr) ) goto 10
    REPEAT THIS UNTIL END OF BOTH STRINGS OR POLY FOUND
    NOW CHECK FOR END OF STRING CONDITION, REQUIRES BACKING UP AND
    USING A PREVIOUS POINT.
    if (endstr) then
       if (blen(1) .eq. 1) then
          string = 2
          other = 1
       else if (blen(2) .eq. 1) then
          string = 1
          other = 2
       else if (model(2,s(string)-1,string) .lt.
                model(2,s(other)-1,other))
   &
      then
   δ
          tmp = other
          other = string
          string= tmp
        endif
       n(string) = s(string) - 1
    endif
    do j=1,3
       verts(j,1) = model(j, s(string), string)
        verts(j,2) = model(j, s(other), other)
        verts(j,3) = model(j, n(string), string)
    enddo
     trarap = interp( verts, reqpt)
     if (trarap .gt. 0) trarap = ( trarap)
998 continue
    return
     end
```

```
Interpolates between 3 points in 3D space.
         C
                        PURPOSE:
         С
                                                       Constructs function of a plane containing the 3 points
         C
                        METHOD:
                                                       and calculates the Z value for the given (X,Y) point.
         C
         C
                                                      CREATED
                                                                                     July 1993
                                                                                                                           H.D.R. Evans
                        HISTORY:
         implicit none
                        save
                                                  gpnts(3, *)
                        real*4
                                                   reqpnt(3)
                        real*4
                                                   v1(3), v2(3)
                        real*4
                                                   rv(3)
                        real*4
                                                    plane(4)
                        real*4
                        real*4
                                                   rv1, rv2, scl
                                                    dotp, disl2
                        real*4
                         integer*4 i
                         Interp = 0.0
                         COMPUTE VECTORS IN PLANE OF THREE POINTS.
         C
ũ
                         do i=1,3
The state of the s
                                v1(i) = gpnts(i, 2) - gpnts(i, 1)
                                v2(i) = gpnts(i, 3) - gpnts(i, 1)
                         enddo
                         DETERMINE NORMAL TO PLANE DEFINED BY V1 AND V2, AND PLANE
                         CONSTANT. PLANE(1) X + PLANE(2) Y + PLANE(3) Z = PLANE(4)
         С
7
                         call CrossP( v1, v2, plane, 3)
plane(4) = DotP(3, GPnts(1,1), 1, plane, 1)
Fu
VALUE WE REQUIRE IS THE Z VALUE AT THE POINT SPECIFIED
С
                         BY THE SOLUTION OF:
                                   Z = (PLANE(4) - PLANE(1)X - PLANE(2)Y) / PLANE(3)
         С
                         IF (Plane(3) .NE. 0) THEN
                                 Interp = (Plane(4) - Plane(1) * ReqPnt(1)
                                                                                       - Plane(2) * ReqPnt(2) ) / Plane(3)
                         else
                                print*, 'plane containing 3 given points is independent of z'
                                print*,'plane = ',plane,char(7)
                                 stop
                         endif
                         return
                         end
                         subroutine crossp(x, y, z, dim)
```

real function interp (gpnts, reqpnt)

```
H.D.R. Evans ESA/ESTEC/WMA
                         July 1993
               CREATED
       HISTORY:
  implicit none
       save
           - 1ST VECTOR
  C
       X
       Y
           - 2ND VECTOR
  C
           - CROSSPRODUCT = X^Y
  C
       DIM - DIMENSION OF X, Y AND Z
  C
       real*4 x(*), y(*), z(*)
       real*4 magz
                 dim
       integer*4
                 i, j, indx
       integer*4
       indx(j) = mod(j + dim - 1, dim) + 1
       do i=1, dim
         magz = 0
         z(i) = x(indx(i+1)) * y(indx(i+2)) -
                x(indx(i+2)) * y(indx(i+1))
       enddo
       return
       end
  real*4 function dotp( n, sx, incx, sy, incy )
                                         ******
  C**********
                Returns the inner (dot) product of SX and SY.
       PURPOSE:
  C
T.
  C
               Basic vector calculations. Vectors must be the
  C
       METHOD:
  С
                same size.
ļ.
  C
                                     H.D.R. Evans ESA/ESTEC/WMA
                         July 1993
       HISTORY: CREATED
  C
3
  1
implicit none
       save
integer*4 n, incx, incy
real*4 sx(*), sy(*)
       integer*4 i, pos
       pos(i,incx) = (i-1)*incx + 1
       dotp = 0
       do i=1,n
         dotp = dotp + sx(pos(i,incx)) * sy(pos(i,incy))
       enddo
       return
       end
   (********************************
       logical function inpoly(a, b, c, pt)
   Returns .TRUE. if the (X,Y) coordinates of point Pt
   С
       PURPOSE:
                 is in the polygon described by the points A,B, and C
   С
   С
   С
       METHOD:
                Pts is decomposed into the sum of the line segments
                 AC and BC, i.e. PTS = a * AC + b * BC. If either a
   С
                 or b is less than zero, then PTS is not subtended by
   C
                 the lines AC & BC.
   С
   C
                 This is then repeated for the AB and CB line segements
   С
```

```
to ensure PTS isn't the other side of the AB line from C.
С
C
                                       H.D.R. Evans ESA/ESTEC/WMA
               CREATED
                          July 1993
C
     HISTORY:
C
implicit none
     save
     real*4 a(2), b(2), c(2), pt(2)
     real*4
              det
     real*4 m (2,2), im(2,2)
     real*4 u(2), v(2), po(2), coeff(2)
     integer*4 i
     do i=1,2! initialise the matrix
        m(1,i) = a(i) - c(i)
        m(2,i) = b(i) - c(i)
        po(i) = pt(i) - c(i)
     enddo
     det = m(1,1) * m(2,2) - m(1,2) * m(2,1)
     if ( det .eq. 0 ) goto 999
     im(1,1) = m(2,2)
     im(2,2) = m(1,1)
     im(1,2) = -m(2,1)
     im(2,1) = -m(1,2)
     coeff(1) = (im(1,1) * po(1) + im(1,2) * po(2))/det
     coeff(2) = (im(2,1) * po(1) + im(2,2) * po(2))/det
     inpoly = ( coeff(1) .ge. 0.0 .and. coeff(2) .ge. 0.0)
     REPEAT THE PREVIOUS, ONLY THIS TIME WITH THE END POINT.
С
     do i=1,2
        m(1,i) = b(i) - a(i)
        m(2,i) = c(i) - a(i)
        po(i) = pt(i) - a(i)
      enddo
      det = m(1,1) * m(2,2) - m(1,2) * m(2,1)
      if ( det .eq. 0 ) goto 999
      im(1,1) = m(2,2)
      im(2,2) = m(1,1)
      im(1,2) = -m(2,1)
      im(2,1) = -m(1,2)
      coeff(1) = (im(1,1) * po(1) + im(1,2) * po(2))/det
      coeff(2) = (im(2,1) * po(1) + im(2,2) * po(2))/det
      inpoly = (coeff(1) .ge. 0.0 .and. coeff(2) .ge. 0.0)
              .and. inpoly
      return
 999 continue
      inpoly = .false.
      print*, char(7)
      stop '***inpoly*** determinant = 0'
            ******************
      real*4 function xinter(x, x1, y1, x2, y2)
 linearly interpolates between (x1,y1) and (x2,y2).
```

```
simple linear interpolation.
  С
      method:
  С
                        july 1993
                                   h.d.r. evans esa/estec/wma
      history:
               created
  С
  implicit none
       save
       real*4 x, x1, x2, y1, y2
       if (x2 .ne. x1) then
         xinter = y1 + (x- x1)*(y2-y1)/(x2-x1)
         xinter = y1
       endif
       return
  integer*4 function estrng( map, header, e, len)
  C****************************
  C
       RETURNS INDEX IN THE AX8 MAP WHERE REQUESTED ENERGY STRING STARTS.
  C
  implicit none
ű
       save
       real*4
              header(*), energy
       real*4
       integer*4 index, len
integer*4 map(*)
       integer*4 epos, escl, maplen
            epos, escl, maplen / 1, 4, 8/
3
           : length of current energy string
ž.å
       epos : offset in the energy string of the energy
maplen: position in the header of the total map length
L
       escl : position in the header of the energy scale factor
.
Lake
index = 1
       energy = 0
       if ((e.le.energy) .or. (index .gt. header(maplen))) goto 20
   10
         len = map( index)
         energy = 1.0 * map( index + epos) / header(escl)
         if ( e .gt. energy) index = index + len
       goto 10
   20
       continue
       estrng = index
       return
  C**********************************
       integer*4 function lstrng( estr, header, 1, len)
  C.
       Returns the index in the Energy string (ESTR)
       that the requested L string starts.
  implicit none
       save
       real*4
       real*4
               header(*)
```

С

```
The state of the s
```

```
real*4
              mapl
     integer*4 index, len
     integer*4 estr(*)
     integer*4 slen, lpos, lscl
                                   1, 5/
           slen, lpos, lscl / 1,
     data
     slen - position in the e string of the e string length
C
     lpos - offset in the l string of the l value
С
     lscl - position in the header of the l scale factor
C
     index = position of the first 1 string in the e string
     index = 3
     mapl = 0
     if ( l .eq. 0 ) then
       lstrng = index - 1
       return
     endif
     if ((l.le.mapl) .or. (index .gt. estr(slen))) goto 20
10
       len = estr( index)
       mapl = estr( index + lpos) / header( lscl)
       if ( l .gt. mapl) index = index + len
     goto 10
20
     continue
     lstrng = index - 1
     return
     end
C****************************
     subroutine evals ( map, header, e, npts)
Searches through the Ax8 model for all of the energies it contains.
implicit none
     save
                 header(*)
     real*4
     real*4
                 e(*)
                 map(*)
     integer*4
     integer*4
                 npts, index
     integer*4 epos, escl, maplen
            epos, escl, maplen / 1, 4, 8/
     data
             : OFFSET IN THE ENERGY STRING OF THE ENERGY
C
     EPOS
             : POSITION IN THE HEADER OF THE ENERGY SCALE FACTOR
C
     ESCL
            : POSITION IN THE HEADER OF THE TOTAL MAP LENGTH
C
     index
            = 1
             = 0
     npts
 10
     continue
        npts = npts + 1
        e(npts) = map( index + 1) / header( escl)
        index = index + map(index)
     if (index .le. header(maplen) .and. map(index) .ne. 0) go to 10
     return
     end
C***********************************
```

```
Searches through Energy string for all of the L values it contains.
  estr(*)
       integer*4
       real*4
                   header(*)
       real*4
                   1(*)
       integer*4
                   npts
       SLEN - POSITION IN THE E STRING OF THE E STRING LENGTH
  C
       LPOS - OFFSET IN THE L STRING OF THE L VALUE
  С
       LSCL - POSITION IN THE HEADER OF THE L SCALE FACTOR
       integer*4 slen, lpos, lscl
              slen, lpos, lscl /
                                1,
                                     1,
                                          5 /
       index
              = 3
              = 0
       npts
       if (index .ge. estr(slen)) goto 20
   10
          npts = npts + 1
          l(npts) = estr( index + lpos) / header( lscl)
          index = index + estr( index)
       goto 10
   20
       continue
return
end
subroutine bb0val(lstr, header, bob0, lnflux, npts)
  Returns the BoBo and Log10(flux) values contained in the L string
C
  C
       (MAP)
  C**********************
       implicit none
₹.
       save
ini.
                   lstr(*)
       integer*4
ĩIJ
       real*4
                   header(*)
real*4
                   bob0(*)
Ŀå
       real*4
                   lnflux(*), i
ű
       integer*4
                   npts
١...
       integer*4
                   len
       integer*4 slen, bscl, flxscl, flxinc, flxoff
       data
              slen, bscl, flxscl, flxinc, flxoff
                      6,
                            7, -256,
               1,
       SLEN - POSITION IN THE B STRING OF THE B STRING LENGTH
  C
  C
       LPOS - OFFSET IN THE L STRING OF THE L VALUE
       LSCL - POSITION IN THE HEADER OF THE L SCALE FACTOR
  C
       FLXSCL - POSITION IN THE HEADER OF THE LNFLUX SCALE FACTOR
  C
  С
       FLXINC - UNSCALED INCREMENT IN B BETWEEN STRING VALUES
  C
       FLXOFF - OFFSET IN B STRING OF THE BO LNFLUX VALUE
       npts = 1
       len = lstr( slen)
       bob0(1) = 1.0
       lnflux(1) = lstr( flxoff) / header(flxscl)
       if (len .lt. 4) return
       npts = 0
       i = 4
       if ((i.gt.len) .or. (lstr(i).le. 0)) goto 20
   10
```

subroutine lvals (estr, header, 1, npts)

```
lnflux(i-2) = (lstr(flxoff) + flxinc*(i-flxoff)) /
                                                         header(flxscl)
                                                 = bob0(i-flxoff) + lstr(i)/header(bscl)
                        bob0(i-2)
                        i = i + 1
                  goto 10
                  continue
        20
                  return
                  end
      subroutine bvals (lstr, header, b, lnflux, npts)
      C**********************************
                Returns B and Log10(flux) values contained in L string (MAP)
      implicit none
                  save
                                              header(*),
                                                                       b(*)
                  real*4
                                              lnflux(*), bob0(40)
                  real*4
                                              1, bo, b0
                  real*4
                  integer*4
                                              lstr(*)
                  integer*4
                                              npts, i
                  integer*4 lpos, lscl
                                    lpos, lscl / 2, 5/
                  data
The state of the s
                  LPOS - POSITION IN L STRING OF THE L VALUE
                  LSCL - POSITION IN THE HEADER OF THE L SCALE FACTOR
                              = lstr(lpos) / header( lscl)
                  call bb0val(lstr, header, bob0, lnflux, npts)
                  bo = b0(1)
<u>l</u>
                  do i = 1, npts
æ
                        b(i) = bob0(i) * bo
if (b(i) .eq. 0) lnflux(i) = 0.0
fil
                  enddo
L
                  return
                  end
      subroutine phival (lstr, header, l, phi, lnflux, npts)
      Returns Hybrid and Log10(flux) values contained in L string (MAP)
      implicit
                                         none
                  save
                  real*4
                                              bm, b0(100), bmax
                  real*4
                                              header(*)
                  real*4
                  real*4
                                              lnflux(*)
                  real*4
                                              phi(*)
                  real*4
                                              xphi
                  integer*4
                                               i
                  integer*4
                                               lstr(*)
                  integer*4
                                              npts
                  call bb0val( lstr, header, b0, lnflux, npts)
                  CONVERT THE B VALUES TO THE HYBRID ONES
      С
                  bm = b0 (npts)
                  do 10 i = 1, npts
                        phi(i) = xphi(b0(i), bm)
```

npts = npts + 1

```
if (phi(i) .lt. 0) lnflux(i) = 0.0
     continue
  10
     return
     end
 real*4 function xphi (bob0, bmax)
 Computes the hybrid magnetic field coordinate=
 С
            (B/B0 - 1)
 С
     XPHI= ASIN( -----
 С
            (Bmax/B0 - 1)
 C
 C
     Where Bmax is the atmospheric cutoff value for B.
     If Bmax = 1, then XPHI = 90. ( Arcsin( 1.0) )
 C
 implicit none
     save
     real*4 bob0, bmax
     real*4 sine
     if (bob0 .gt. bmax) then
       ARCSIN( >1.0) --- BOBO BEYOND THE ATMOSPHERIC CUT OFF.
 C
       xphi = -1.0
       return
     endif
     if (bmax .ne. 1) then
       sine = (bob0 - 1)/(bmax-1)
       if ( (-1.le.sine).and.(sine .le.1.0) ) then
         xphi = asin(sine) * 180.0 / 3.1415927
       else
High
         xphi = -1.0
       endif
     else
xphi = -1.0
     endif
return
TU
 ļ.
     real*4 function b0(1)
computes magnetic field strength for an L shell at magnetic equator.
 implicit none
     save
     real*4 l
     if ( 1 .gt. 0 ) then
       b0 = 0.311653 / (1*1*1) ! ????ASK TONY: SHOULWE USE GMAGMO?
     else
       b0 = 0
     endif
     return
     end
```

```
PROGRAM Trapped Proton Driver
C
        IMPLICIT NONE
        SAVE !from original DRIVER supplied by Colborn & Armstrong
      integer*4 ie, iemax, ifile, igo, imod, ne, ns, Mpts
      parameter (iemax = 30) ! MAX NO. OF ENERGIES ALLOWED
      real Evals(iemax)
        INTEGER NLvals
        PARAMETER (NLvals=10)
                         !program now calculates this from period and Norbits.
        INTEGER Ndays
        INTEGER Norbits, Norbsteps
                                    !arguments to subroutine calls, 11-26-97.
        REAL OrbPrecTime !for future use, 11-26-97.
        REAL TrappedFlux(IEmax, NLvals), XLbounds(NLvals)
        REAL RelDwellTime (NLvals)
        REAL Year
        REAL DiffTrappedFlux(IEmax, NLvals)
        INTEGER MARR
        PARAMETER (MARR=5000)
        REAL Eout (MARR), Fluxout (MARR, NLvals)
        REAL OrbIncl, Apogee, Perigee, AscNodeLong, AscNodeDisp, PerigDisp
        INTEGER ILbins
        CHARACTER*80 TrappedFile
        INTEGER Program Code
        DATA Program Code/1/
        INTEGER IpreCalc
        LOGICAL DistBelt, PreCalcFlux
        REAL ELOWER, EUPPER
        DATA ELOWER, EUPPER/1.0E-01, 1.0E+05/
        REAL OrbPeriod
C----
                           CALL TrappedDriverInput (Evals, NE, OrbIncl, Apogee, Perigee,
     #
               AscNodeLong, AscNodeDisp, PerigDisp, TrappedFile, Year,
     #
               XLbounds, ILbins, imod, DistBelt, PreCalcFlux, IPreCalc,
     #
               Ndays, Norbits, Norbsteps)
        CALL Trapped ORBINT (OrbIncl, Apogee, Perigee, AscNodeLong,
     #
               AscNodeDisp, PerigDisp, Evals, NE, TrappedFlux,
     #
               DiffTrappedFlux, Year, XLbounds, ILbins, imod, RelDwellTime,
     #
               DistBelt, PreCalcFlux, IPreCalc,
     #
               Ndays, Norbits, Norbsteps, OrbPeriod, OrbPrecTime)
        CALL Trapped Spectra (Evals, NE, ELOWER, EUPPER, Mpts, ILbins,
     #
               TrappedFlux, DiffTrappedFlux, Eout, Fluxout)
        CALL OutputTrappedFlux(Evals, NE, TrappedFlux, DiffTrappedFlux,
```

```
#
                 imod, RelDwellTime, DistBelt, PreCalcFlux, IPreCalc,
       #
                 ELOWER, EUPPER, Mpts, Eout, Fluxout,
                 Ndays, Norbits, Norbsteps, OrbPeriod, OrbPrecTime)
          STOP
          END
   C**********************************
        subroutine differ(ne, e, fa, fb, d, Ilbins)
   implicit none
        save
        real*4 da, db
        integer*4 i, j, ne, ILbins, L
        INTEGER iemax, NL vals
        PARAMETER (iemax = 30, NLvals=10)
  C
        make array fixed size, so that don't have difficulties with 2D
  С
        array allignment.
        REAL*4 e(iemax),fa(iemax,NLvals)
        REAL*4 fb(iemax, NLvals), d(iemax, NLvals)
101
        routine finds d(i) the differential flux at energy e(i)
        assuming that the spectrum is best represented by an exponential
  C
        ne is number of energies
  С
        fa(i) is integral flux above e(i)
.
Posta
  С
        fa(ne) is defined by routine
e
Lak
        fb(i) is the flux between e(i+1) and e(i)
IJ
la la
        do L = 1, ILbins
        do i = ne, 1, -1
           if (fa(i,L) .gt. 0) go to 2
           d(i,L) = 1 .0e-37
        enddo
        GOTO 99
        if (i \cdot eq. ne) i = i-1
        fa(i+1,L) = fa(i,L) - fb(i,L)
        if (fa(i+1,L) .ne. 0) go to 6
        i = i-1
        if (i .gt. 1) go to 6
        d(1,L) = fa(1,L)/(e(2)-e(1))
        GOTO 99
   6
        db = -alog(fa(2,L)/fa(1,L))/(e(2)-e(1))*fa(1,L)
        do j = 1, i
           da = -alog(fa(j+1,L)/fa(j,L))/(e(j+1)-e(j))*fa(j,L)
  C
           Added error checking on da*db, 11-24-97.
           IF (da*db .GE . 0.0) THEN !da*db should be >= 0 for physical solutions
             d(j,L) = sqrt(da*db)
```

TrappedFile, OrbIncl, Apogee, Perigee, AscNodeLong,

AscNodeDisp, PerigDisp, Year, XLbounds, ILbins, PROGRAM CODE,

#

ELSE

```
the fact that the fact that the first the fact that the fa
```

```
d(j,L) = 0.0
         ENDIF
         db = da*fa(j+1,L)/fa(j,L)
      enddo
      d(i+1,L) = da*fa(i+1,L)/fa(i,L)
                                       ! for protons
99
      CONTINUE !for going to next L-value bin instead of return
      enddo !stepping through L-bins.
      return
      end
SUBROUTINE TrappedDriverInput(Evals, NE, OrbIncl, Apogee, Perigee,
     #
                      AscNodeLong, AscNodeDisp, PerigDisp, TrappedFile,
     #
                      Year, XLbounds, ILbinsum, imod,
     #
                      DistBelt, PreCalcFlux, IPreCalc,
     #
                      Ndays, Norbits, Norbsteps, OrbPrecTime)
        IMPLICIT NONE
        REAL OrbIncl, Apogee, Perigee, AscNodeLong, AscNodeDisp, PerigDisp
C
        Note that the eccentricity is calculated here to decide if
        need to read PerigDisp. The eccentricity is also recalculated
C
C
        in the initialization CALL ORBIT(1,...) case.
С
C
        This makes the input driver independent of the actual computational
        routines, so that it will be easier to modify and interface with other
        space environment routines.
        REAL E, Re
                      !eccentricity and radius of Earth
        PARAMETER (Re=6371.2)
        REAL ApPerSwitch
        INTEGER IERR, IACCEPT
        DATA IERR/0/
        INTEGER Ndays
        INTEGER Norbits, Norbsteps
                                   !now passed as arguments, 11-26-97.
        REAL OrbPrecTime !for future use, 11-26-97.
        CHARACTER*80 TrappedFile
        INTEGER NLvals, I, L, ILbinMax, ILbinsum, imod
        PARAMETER (NLvals=10)
        REAL XLbounds (NLvals), XLinfinite, Year, YearMin, YearMax
        PARAMETER (XLinfinite=1.0E+06)
C
        Appropriate epochs for AP8MIN & AP8MAX L-value calculations
        PARAMETER (Yearmin=1964.0, Yearmax=1970.0)
       REAL XLdummy
      INTEGER iemax
     parameter (iemax = 30)
                              ! MAX NO. OF ENERGIES ALLOWED
     real evals(iemax)
      integer ne
```

```
the first of the state of the s
```

```
define energy grid for test case
C
      energy values often used in space station calculation
C
С
     These are hardwired into calculation for now.
C
     Changed E to Evals (since E is the now the eccentricity)
       INTEGER IpreCalc, IFLUXtype
       LOGICAL DistBelt, PreCalcFlux
       INTEGER Idummy !For diagnostic message on number of I/O parameters.
C-----
       NE = 29
       EVALS(1) = 1.0
       EVALS(2) = 2.0
       EVALS(3) = 3.0
       EVALS(4) = 4.5
       EVALS(5) = 6.
       EVALS(6) = 8.
       EVALS(7) = 10.
       EVALS(8) = 15.
       EVALS(9) = 20.
       EVALS(10) = 30.
       EVALS(11) = 50.
       EVALS(12) = 80.
       EVALS(13) = 100.
       EVALS(14) = 150.
       EVALS(15) = 200.
       EVALS (16) = 250.
       EVALS(17) = 300.
       EVALS(18) = 350.
       EVALS(19) = 400.
       EVALS (20) = 450.
       EVALS(21) = 500.
       EVALS(22) = 550.
       EVALS(23) = 600.
       EVALS (24) = 800.
       EVALS(25) = 1000.
       EVALS (26) = 3000.
       EVALS(27) = 1.0E + 04
       EVALS(28) = 3.0E + 04
       EVALS(29) = 1.0E + 05
      WRITE(6,1000)
      WRITE (6, 1001)
С
       initialize boundaries L-value bins
       XLbounds(1) = 0.0
       DO L=2, NL vals
         XLbounds(L)=XLinfinite
       ENDDO
 9390
       CONTINUE
       CALL RETRY INPUT (IERR)
       WRITE(*,390)
       IFLUXtype=0
       READ(*,*,ERR=9390,IOSTAT=IERR)IFLUXtype !installed, 11-26-97.
       PreCalcFlux=.FALSE. !initialize not to use pre-calculated Flux
```

```
!hardwire off for now.
           DistBelt=.FALSE.
           IpreCalc=0
           IF (IFLUXtype .NE. 0) THEN
    9391
             CONTINUE
             CALL RETRY INPUT (IERR)
             WRITE(*,391)
             READ(*,*,ERR=9391,IOSTAT=IERR)IpreCalc
             PreCalcFlux=.TRUE.
                                !hardwire off for now, since we do not include a
             DistBelt=.FALSE.
                                 !"disturbed belt" model
   C
             Use quiet-time, 51.6 degrees as the default case.
             IF (IpreCalc .LT. 0 .OR. IpreCalc .GT. 3) IpreCalc=0
             CONTINUE
    9427
             CALL RETRY INPUT (IERR)
             WRITE(*,427)
             READ(*,428,ERR=9427,IOSTAT=IERR)TrappedFile
             CALL CHECK_OUTPUT_FILE(TrappedFile, IACCEPT)
             IF (IACCEPT.NE.0) GOTO 9427
□ C
             The SUBROUTINE OutputTrappedFlux presently takes these variables
             from the header information in the pre-calculated files.
  C
             IMOD is also taken from that header information. Only IpreCalc
             is re-checked against that header information. 12-1-97.
  C
  C
  CF?
                IF (IpreCalc .EQ. 0 .OR. IpreCalc .EQ. 1) THEN
                  OrbIncl=51.6
  CF?
  CF?
                  Apogee=400.0
  CF?
                  Perigee=400.0
  CF?
                ELSEIF (IpreCalc .EQ. 2 .OR. IpreCalc .EQ. 3) THEN
  CF?
                  OrbIncl=28.5
  CF?
                  Apogee=450.0
  CF?
                  Perigee=450.0
  CF?
                ENDIF
<sup>to</sup>al C
             The pre-calculated fluxes are not available (and not divided into
  С
             L-bins)
             ILbinsum=1
             RETURN
           ENDIF
           Moved the questions and answers on AP8MIN or AP8MAX here, 12-11-97.
    9454
           CONTINUE
           CALL RETRY INPUT (IERR)
           WRITE (*, 454)
           READ (*, *, ERR=9454, IOSTAT=IERR) imod
           IF (imod .LT. 0 .OR. imod .GT. 1) imod=0
         imod = imod + 1
                           Ţ
                               model no. for subroutine blccoords
                               model no. for subroutine trapped protons
           IF (imod .EQ. 1) YEAR=1964.0
           IF (imod .EQ. 2) YEAR=1970.0
```

T

200

!hardwired off for now.

```
С
                              End of moved questions & answers on AP8MIN or AP8MAX, 12-11-97.
        C
                              What is the altitude at apogee?
        C
           9420
                              CONTINUE
                              CALL RETRY INPUT (IERR)
                              WRITE(*,420)
                              READ (*, *, ERR=9420, IOSTAT=IERR) Apogee
        C
        C
                              WHAT IS THE ALTITUDE AT PERIGEE?
        С
                             CONTINUE
           9400
                              CALL RETRY INPUT (IERR)
                              WRITE(*,400)
                              READ (*,*,ERR=9400,IOSTAT=IERR) Perigee
        С
                             allow the user to specify apogee and perigee in either order
                              instead of performing unintended calculation which sets eccentricity
        C
        C
                              to zero and using Perigee variable (actual apogee) to produce
                              a circular orbital altitude in ORBIT routine.
        С
                              IF (Perigee .GT. Apogee) THEN
                                      ApPerSwitch=Apogee
                                      Apogee=Perigee
                                      Perigee=ApPerSwitch
The line and the line and the same of the 
                                      WRITE(*,430)
                             ENDIF
                              E= (Apogee-Perigee) / (Apogee+Perigee+2.*Re)
                              IF (E.LT..00001) E=0.
        С
                              WHAT IS THE ORBITAL INCLINATION?
į.
          9405
                             CONTINUE
CALL RETRY INPUT (IERR)
L
                              WRITE(*,405)
la.
                             READ (*, *, ERR=9405, IOSTAT=IERR) OrbIncl
T C
                             READ in number of orbits, 11-26-97.
        C
                             Have Removed "FAST" option, i.e. must enter Ascending Node information
        C
        С
                              Retain these initializations in case want to hardwire ascending
        C
                             node information at future time.
                             AscNodeLong=0.
                             AscNodeDisp=0.
                             PerigDisp=0.
        C
                             Modified WRITE statement, 12-11-97.
                             IF (E.NE.O.) THEN
                                   Idummy=3
                             ELSE
                                   Idummy=2
                             ENDIF
                              WRITE(*,409)Idummy
```

ű.

```
С
           WHAT IS THE INITIAL LONGITUDE OF THE ASCENDING NODE?
   С
    9410
           CONTINUE
           CALL RETRY INPUT (IERR)
           WRITE(*,410)
           READ (*, *, ERR=9410, IOSTAT=IERR) AscNodeLong
   C
   С
           WHAT IS THE INITIAL DISPLACEMENT FROM THE ASCENDING NODE?
   C
           CONTINUE
    9415
           CALL RETRY INPUT (IERR)
           WRITE(*,415)
           READ (*, *, ERR=9415, IOSTAT=IERR) AscNodeDisp
           IF (E.NE.O.) THEN
                                    !Only read in XI if eccentricity is nonzero
   С
   С
               What is the displacement of the perigee from the ascending node?
   C
            CONTINUE
    9425
            CALL RETRY INPUT (IERR)
            WRITE(*,425)
            READ (*, *, ERR=9425, IOSTAT=IERR) PerigDisp
           ENDIF
IF ( (AscNodeLong .NE. 0.0) .OR. (AscNodeDisp .NE. 0.0) .OR.
                (PerigDisp .NE. 0.0) ) WRITE(*,426)
           Moved the entry of Norbits here, 12-11-97.
           CONTINUE
    9408
           CALL RETRY INPUT (IERR)
1
           WRITE (*, 408)
   CF?
              READ (*, *, ERR=9408, IOSTAT=IERR) Norbits, Norbsteps
L
           READ (*, *, ERR=9408, IOSTAT=IERR) Norbits
Norbsteps=200
                             !hardwired at 200 for now, 11-26-97.
           IF (Norbits .LE. 0) Norbits=200
  CF?
              IF (Norbsteps .LE. 0) Norbsteps=200
   C
           End of moved section for entry of Norbits, 12-11-97.
    9450
           CONTINUE
           CALL RETRY_INPUT(IERR)
           WRITE (*, 450)
           READ (*, *, ERR=9450, IOSTAT=IERR) ILbinMax
           IF (ILbinMax .LT. 0) ILbinMax=0
           IF (ILbinMax .GT. NLvals) THEN
              WRITE(*, 456)
              ILbinMax = NLvals
           ENDIF
           IF (ILbinMax .GT. 0) THEN
             WRITE(*,451) ILbinMax
    9451
               CONTINUE
               CALL RETRY INPUT (IERR)
               READ (*, *, ERR=9451, IOSTAT=IERR) (XLbounds(L), L=1, IlbinMax)
```

#

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IF (ILbinMax .EQ. 1 .AND. XLBOUNDS(1) .EQ. 0.0) WRITE(*,458)
        ENDIF
        IF (XLbounds(1) .LT. 0.0) XLbounds(1)=0.0
C
        Start DO loop at 1, so that ILbinMax=2 will be properly handled
        This SUBROUTINE insists the L-values are in increasing order.
C
        If this is not the case, all subsequent L-value bins will be
C
C
        ignored.
        DO L=1, ILbinMax
          IF (XLbounds(L) .LT. XLbounds(1)) THEN
              WRITE(*,452) XLbounds(L),XLinfinite
              XLbounds (L) = XLinfinite
          ENDIF
          IF (L .GE. 2) THEN
            IF (XLbounds(L) .LE. XLbounds(L-1)) THEN
              WRITE(*,452) XLbounds(L), XLinfinite
              XLbounds (L) = XLinfinite
            ENDIF
          ENDIF
        ENDDO
        ILbinsum=1
       DO L=1, ILbinMax
          IF ( (L .GE. 2) .AND. (XLbounds(L) .LT. XLinfinite) )
                ILbinsum=ILbinsum+1
        ENDDO
        IF (ILbinMax .NE. ILbinsum .AND. ILbinMax .NE. 0) THEN
          WRITE(*,453)ILbinMax,ILbinsum
          ILbinMax=ILbinsum
        ENDIF
9428
       CONTINUE
        CALL RETRY INPUT (IERR)
        IF (ILbinMax .EQ. 0 .OR. (ILbinMax .EQ. 1 .AND.
           XLBOUNDS(1) .EQ. 0.0) ) THEN
         WRITE(*,427)
         READ(*,428,ERR=9428,IOSTAT=IERR)TrappedFile
          CALL CHECK OUTPUT FILE (Trappedfile, IACCEPT)
          IF (IACCEPT.NE.0) GOTO 9428
        ELSE
          WRITE(*,455)ILbinMax
         READ(*,428,ERR=9428,IOSTAT=IERR)TrappedFile
         CALL CHECK OUTPUT FILE (Trappedfile, IACCEPT)
          IF (IACCEPT.NE.0) GOTO 9428
         DO I=1, LEN (TrappedFile)
            IF (TrappedFile(I:I) .EQ. '.') THEN
                TrappedFile=TrappedFile(1:I-1)
           ENDIF
          ENDDO
       ENDIF
       RETURN
      FORMAT(1X,'Orbit averaging using ESA AP8 Models.',/)
     FORMAT(' This program will calculate the omnidirectional',
1001
```

```
' trapped proton flux',
               /,' to a',
                ' spacecraft orbiting inside the magnetosphere. The',
               /,' calculated trapped flux is for demonstration purposes',
                ' only, and is NOT part of the standard CREME96 software.',
        &//,' NOTE: Before running this or any other CREME96 programs',
        & ' please define three ',
        & /,' logicals: ',/,
        & /,4x,' CREME96
                               as the directory where CREME96 source',
                 ' & executables reside.',
        & /,4x,' CR96TABLES
                             as the directory in which CREME96 data',
                ' tables reside.',
        & /,4x,' USER
                               as the directory in which output files',
                ' should be written.',
        & //,' Now begin specification of the ESA AP8 calculation: ',/)
    390
          FORMAT(1X,'Enter 0 to directly calculate the trapped',
                      'proton fluxes, or',/,1X,'Enter 1 to test the ',
        &
                      'pre-calculated trapped proton interface.',/,3X,
                     '[NOTE: The pre-calculated fluxes are presently ',
        &
                     'test case results which', /, 11X,
                     'are intended only as a software test.]')
    391
          FORMAT(1X, 'Enter 0 for Space Station (51.6 deg., 400 km)',
              ' orbit (ISSA), ESA-AP8MAX,',
        & /,7X,'1 for ISSA, ESA-AP8MIN,',
& /,7X,'2 for 28.5 deg. (450 km), ESA-AP8MAX,',
& /,7X,'3 for 28.5 deg. (450 km), ESA-AP8MIN.')
    420
          FORMAT(/,1X,'Enter altitude at apogee (kilometers): ')
    400
          FORMAT(/,1X,'Enter altitude at perigee (kilometers): ')
    430
          FORMAT(/,1X,'Input apogee < perigee, have been interchanged.')
2 5
æ
    405
          FORMAT(/,1X,'Enter orbital inclination (degrees): ')
ļ.
TU c
          Modified 12-11-97.
FORMAT(/,1X,'The following ',I1,' input parameters are ',
   409
1
            'most relevant to situations in which',
       &
       & /,1X,'the actual orbital path is known',
1
              ' or in which mission critical operations',
ĮĮ.
       & /,1x,' are planned.',
        & //,3X,' [Recommended values are 0.0, unless you wish to examine',
        & /,3X,'a very specific orbital segment.]')
          FORMAT(/,1X,'Enter initial longitude of ascending node',
    410
                   1X, '[Recommended = 0.0 (degrees)]:')
         FORMAT(/,1X,'Enter initial displacement from ascending',
    415
                 ' node',1X,'[Recommended = 0.0 (degrees)]:')
          FORMAT(/,1X,'Enter displacement of perigee from',
       1 'ascending node',1X,'[Recommended = 0.0 (degrees)]:')
    426
          FORMAT(/,1X,'Note: for studies sensitive to a specific',
       & 'orbital segment, you should be',
       & /,1X,'aware that the trapped proton',
       & ' calculations are averaged over 7 days at present. This',
       & /,1X,'parameter can be easily reset by modifying',
          ' the TRAPPED_ORBINT subroutine, but is ',
       & /,1x,'not provided as a general-use input parameter.')
```

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```
FORMAT(/,1X,'Enter name of output trapped proton file:',
427
              1,'
                     [Recommended: something.TRP]')
    &
       FORMAT (A80)
428
       FORMAT(/,1X,'Enter the number of desired TRP L-value bins ',
450
            '(1 - 10):',
    33
    & /,3X,'[Recommended default = 0, i.e.',
            'one trapped flux calculation for the entire orbit.]')
451
       FORMAT (/, 1X,
            'Enter the lower limits of the ', I2,' L-value bins: ',
    &
    & /,3X,'[A typical scenario could be to request 4 bins as the',
            ' as the previous entry.',
    &
    & /,3X,'Then, entries of 0.0, 2.0, 4.0, and 6.0',
            ' would subdivide the orbit into',
    & /,3x,'sections with L < 2, L = 2-4, L = 4-6, and L > 6.]',
    & //,1X,'NOTE: The L-value is a magnetic coordinate roughly',
           ' corresponding to the',
    & /,1X,'distance in Earth Radii to the',
            ' magnetic field line at the magnetic equator.',
    &
    & /,1x,'For example, a geosynchronous orbit is roughly L = 6.6,',
            ' the geographic equator',
    & /,1x,'is about L = 1, and the heart of the',
          ' South Atlantic Anomaly (SAA) is roughly at',
    \& /,1x,'L = 1.2 - 2.',
    & ' Calculated L-values slightly less than 1 do occur; using',
    & /,1X,'a lower limit of L = 0 will account for these.')
       FORMAT(1X,'The L-values MUST be entered in increasing order',
452
    & /,1X,'the L-value of ',F10.2,' has been reset to ',F10.2)
       FORMAT(1X,'The number of L-values bins has been reset',
    & /,1X,' from ',I2,' to ',I2)
      FORMAT(/,1X,
 454
             'Enter 0 (default) for AP8MIN (1964)',1X,
    &
             'or 1 for AP8MAX (1970).')
    &
 455
      FORMAT(/,1X,'Enter root name of output TRP files:',
    & /,1X,'[NOTE: There will be ',I2,' output files, and',
          ' the files for the different L-value'
     & /,1x,' bins will',
             ' be called something.TR# (# = 1,2,...,9,X)]')
 456
       FORMAT(1X,'Only 10 L-values are allowed.')
       FORMAT(1X, 'Calculation reset to whole orbit option, since',
 458
               1X, 'choosing 1 L bin',
    &
             /,1X,' with a minimum L-value equal to 0 is equivalent to',
     &
     &
               1X,'the entire orbit.')
 408
       FORMAT(/,1X,
             'Enter Number of orbits to integrate (default = 200)')
    δ.
CF? 408
          FORMAT(/,1X,
CF?
        &
               'Enter Number of orbits to integrate (default = 200)',/,
CF?
        &
                3X, 'and Number of steps per orbit')
              !TrappedDriverInput routine
        END
```

```
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```
SUBROUTINE OutputTrappedFlux(Evals, NE, TrappedFlux,
               DiffTrappedFlux, TrappedFile, OrbIncl,
     #
               Apogee, Perigee, AscNodeLong, AscNodeDisp, PerigDisp,
     #
               Year, XLbounds, ILbins, PROGRAM_CODE, imod, RelDwellTime,
     #
               DistBelt, PreCalcFlux, IPreCalc, ELOWER, EUPPER, MPTS,
               Eout, Fluxout, Ndays, Norbits, Norbsteps,
     #
               OrbPeriod, OrbPrecTime)
C
      TEST Routine for writing the orbit-averaged AP8 trapped proton fluxes.
С
C
      IMPLICIT NONE
      INTEGER*4 I, L, OUTUNIT, IOLEN, NE
      DATA OUTUNIT/2/
      INTEGER*4 NHEADER
      CHARACTER*80 TrappedFile, TEMPFILE
      CHARACTER*9 CREATION DATE
      CHARACTER*8 CREATION_TIME
      INTEGER iemax, NL vals, ILbins
      PARAMETER (iemax=30, NLvals=10)
      REAL*4 YEAR
      INTEGER*4 VERSION_NUMBER, PROGRAM_CODE, imod
      REAL*4 ELOWER, EUPPER
      CHARACTER*12 TARGET
      DATA TARGET/'UNSHIELDED '/
      REAL TrappedFlux(iemax, NLvals), Evals(iemax), XLbounds(NLvals)
      REAL RelDwellTime(NLvals)
      REAL DiffTrappedFlux(iemax, NLvals)
      INTEGER MARR, K, MPTS
      PARAMETER (MARR=5000)
      REAL Eout(MARR),Fluxout(MARR,NLvals)
      REAL OrbIncl, Apoque, Perigee, AscNodeLong, AscNodeDisp, PerigDisp
      REAL XLinfinite
      PARAMETER (XLinfinite=1.0E+06)
      REAL Fconv
CC
       For converting flux to /sr-m**2-s from /cm**2-day, assuming isotropic
              TrappedFlux is in /sr-m**2-s, DiffTrappedFlux in /sr-m**2-s-MeV
CC
       flux.
       Fconv = 1.0E+4/(4*PI*86400).
CC
CC
        PARAMETER (Fconv=9.210356E-02)
      PARAMETER (Fconv=7.957747155E+02) ! 86400 * other Fconv (commented out)
      CHARACTER*4 FEXT(10)
      DATA FEXT/'.TR1','.TR2','.TR3','.TR4','.TR5','.TR6','.TR7',
```

```
'.TR8','.TR9','.TRX'/
```

INTEGER IpreCalc
LOGICAL DistBelt, PreCalcFlux

INTEGER CREME96_OPEN, stat

INTEGER Ndays, Norbits, Norbsteps !passed as arguments, 11-26-97.

C Add orbital period, 12-1-97. OrbPrecTime is for future use, 11-26-97. REAL Orbperiod,OrbperiodHrs,OrbPrecTime

C Note that IDistBeltOutput is presently neither used nor output, 11-26-97.

INTEGER IPreCalcOutput, IDistBeltOutput !11-26-97.

INTEGER IPreCalcTmp !12-1-97, for internal consistency & error checking.

INTEGER INPUNIT !11-26-97 DATA INPUNIT/3/

REAL PCfluxpts(iemax), DiffPCfluxpts(iemax) !11-26-97
INTEGER VERSION_TMP, CODE_TMP !11-26-97

REAL PCFluxes (MARR) !11-26-97

INTEGER IZLOW, IZHIGH !11-26-97

CHARACTER*6 APtitle

CHARACTER*80 PreCalcFile, TITLELINE

C-----

CALL GET_CREME96_VERSION(VERSION_NUMBER)
IPreCalcOutput=0 !11-26-97
OrbperiodHrs=Orbperiod/3600.0

IF (PreCalcFlux) THEN

DistBelt=.FALSE. !Local variable for header output file.

IPreCalcOutput=1 !Local variable for header output file.

cf? IpreCalc = 1 & 3 are AP8MIN calculations, 11-26-97.

IF ((IpreCalc .EQ. 1) .OR. (IpreCalc .EQ. 3))

cf? & DistBelt=.TRUE.

ENDIF

1

3

Just the

cf?

C

IDistBeltOutput=0

cf? IF (DistBelt) IDistBeltOutput=1

IF (.NOT. PreCalcFlux) THEN !11-27-97.

C Open output file and write header

C ILbins = 0 & ILbins = 1 from input routine are treated

C as ILbins = 1 for output, since they are stored in the

C same location in the array.

IF (ILbins .EQ. 1 .AND. XLBOUNDS(1) .EQ. 0.0) THEN
 OPEN(UNIT=OUTUNIT,STATUS='NEW',FILE='USER:'//TrappedFile)
 stat = creme96_open(TrappedFile,'user',outunit,'new')

CALL DATE (CREATION DATE)

```
CALL TIME (CREATION_TIME)
  C
        Now prepare header for output file:
  C
          NHEADER=23
          WRITE (OUTUNIT, 990) NHEADER, TrappedFile (1:70),
                 VERSION NUMBER, PROGRAM CODE
       Se
          WRITE(OUTUNIT, 992) VERSION_NUMBER, CREATION_DATE, CREATION_TIME
          WRITE(OUTUNIT,404) OrbIncl,Apogee,Perigee,AscNodeLong,
                         AscNodeDisp, PerigDisp
       #
          RelDwellTime should be 1.0 in this case.
  С
          IF (imod .EQ. 1)
             WRITE(OUTUNIT,405) 'AP8MIN',imod-1,IPreCalcOutput,
                         RelDwellTime(Ilbins), XLbounds(ILbins),
       #
                         XLbounds(Ilbins+1)
        #
           IF (imod .EQ. 2)
             WRITE(OUTUNIT,405) 'AP8MAX',imod-1,IPreCalcOutput,
        #
                         RelDwellTime(Ilbins), XLbounds(ILbins),
        #
                         XLbounds(Ilbins+1)
cf?
           WRITE(OUTUNIT,411) Norbits, Norbsteps
              WRITE(OUTUNIT,411) Norbits, Norbsteps, OrbPrecTime
           WRITE(OUTUNIT,412) OrbPeriodHrs
WRITE (OUTUNIT, 9195)
يَّعَيْنَ
           WRITE (OUTUNIT, 9196)
           DO I=1, NE, 2
               WRITE(OUTUNIT, 9200) (Evals(K), TrappedFlux(K, ILbins) *Fconv,
Jan.
               DiffTrappedFlux(K,ILbins)*Fconv,K=I,I+1)
£,£
           END DO
L.
           WRITE (OUTUNIT, 9100) ELOWER, EUPPER, MPTS, 1, 1, TARGET, YEAR, NE,
                 VERSION_NUMBER, PROGRAM CODE
           WRITE (OUTUNIT, 100)
           Write trapped proton fluxes to file in standard CREME96 format.
   C
           WRITE(OUTUNIT, 200) (Fconv*FluxOut(K, ILbins), K=1, Mpts)
           WRITE (OUTUNIT, 100)
           CLOSE (OUTUNIT)
         ELSE
           DO I=1,66
              IF (TrappedFile(I:I) .NE. ' ' .AND.
                  TrappedFILE(I+1:I+1) .EQ. '') IOLEN=I
           ENDDO
           DO L=1, ILbins
                TEMPFILE=TrappedFILE(1:IOLEN)//FEXT(L)
                 OPEN (UNIT=OUTUNIT, STATUS='NEW', FILE='USER: '//TEMPFILE)
```

```
stat = creme96_open(TEMPFILE,'user',outunit,'new')
                CALL DATE (CREATION DATE)
                CALL TIME (CREATION TIME)
                Now prepare header for output file:
  C
                NHEADER=23
                WRITE (OUTUNIT, 990) NHEADER, TEMPFILE (1:70),
                       VERSION NUMBER, PROGRAM_CODE
        &
                WRITE (OUTUNIT, 992) VERSION_NUMBER, CREATION_DATE,
                       CREATION TIME
        &
                WRITE(OUTUNIT, 404) Orbincl, Apogee, Perigee, AscNodeLong,
                                AscNodeDisp, PerigDisp
        #
                IF (L .LT. NLvals) THEN
                   IF (imod .EQ. 1)
                     WRITE(OUTUNIT,405) 'AP8MIN',imod-1,IPreCalcOutput,
        #
                            RelDwellTime(L), XLbounds(L), XLbounds(L+1)
        #
                   IF (imod .EQ. 2)
                     WRITE(OUTUNIT, 405) 'AP8MAX', imod-1, IPreCalcOutput,
        #
                            RelDwellTime(L), XLbounds(L), XLbounds(L+1)
         #
the that the first that the that the
                ELSE
                   IF (imod .EQ. 1)
                     WRITE(OUTUNIT,405) 'AP8MIN',imod-1,IPreCalcOutput,
         #
                            RelDwellTime(L), XLbounds(L), XLinfinite
                   IF (imod .EQ. 2)
                     WRITE(OUTUNIT, 405) 'AP8MAX', imod-1, IPreCalcOutput,
                            RelDwellTime(L),XLbounds(L),XLinfinite
                 ENDIF
1
              WRITE(OUTUNIT,411) Norbits, Norbsteps
                  WRITE(OUTUNIT, 411) Norbits, Norbsteps, OrbPrecTime
   cf?
L.
            WRITE (OUTUNIT, 412) OrbPeriodHrs
WRITE (OUTUNIT, 9195)
              WRITE (OUTUNIT, 9196)
              DO I=1, NE, 2
                 {\tt WRITE} \, ({\tt OUTUNIT}, {\tt 9200}) \quad ({\tt Evals} \, ({\tt K}) \, , {\tt TrappedFlux} \, ({\tt K}, {\tt L}) \, \star {\tt Fconv} \, ,
                 DiffTrappedFlux(K,L)*Fconv,K=I,I+1)
         #
               END DO
            WRITE(OUTUNIT, 9100) ELOWER, EUPPER, MPTS, 1, 1, TARGET, YEAR, NE,
                   VERSION NUMBER, PROGRAM_CODE
               WRITE (OUTUNIT, 100)
               Write trapped proton fluxes to file in standard CREME96 format.
   С
               WRITE (OUTUNIT, 200) (Fconv*FluxOut(K, L), K=1, Mpts)
               WRITE (OUTUNIT, 100)
               CLOSE (OUTUNIT)
                           !number of L-bins.
             ENDDO
```

!choosing between whole orbit and L-bin options.

ENDIF

```
!handle pre-calculated trapped fluxes, 11-26-97.
      ELSE
        Presently, there are no L-bins for pre-calculated trapped fluxes.
С
        IF (IPreCalc .EQ. 0) PreCalcFile='IPREC0.TRP'
        IF (IPreCalc .EQ. 1) PreCalcFile='IPREC1.TRP'
        IF (IPreCalc .EQ. 2) PreCalcFile='IPREC2.TRP'
        IF (IPreCalc .EQ. 3) PreCalcFile='IPREC3.TRP'
        stat = creme96_open(PreCalcFile,'cr96tables',inpunit,'old')
        stat = creme96_open(TrappedFile,'user',outunit,'new')
        CALL DATE (CREATION_DATE)
        CALL TIME (CREATION_TIME)
        NHEADER=23
        WRITE (OUTUNIT, 990) NHEADER, TrappedFile(1:70),
              VERSION_NUMBER, PROGRAM CODE
     &
        WRITE(OUTUNIT, 992) VERSION_NUMBER, CREATION_DATE, CREATION_TIME
                                        !dummy title lines from creation of
        READ(INPUNIT, 9194)TITLELINE
                                        !pre-calculated trapped fluxes.
        READ(INPUNIT, 9194)TITLELINE
        READ(INPUNIT, 1404) OrbIncl, Apogee, Perigee, AscNodeLong,
                       AscNodeDisp, PerigDisp
     #
        WRITE (OUTUNIT, 404) OrbIncl, Apogee, Perigee, AscNodeLong,
                       AscNodeDisp, PerigDisp
     #
        READ(INPUNIT, 1405) APtitle, imod, IPreCalcTmp
         READ(INPUNIT, 1406)RelDwellTime(Ilbins),XLbounds(ILbins),
                       XLbounds (Ilbins+1)
     #
         WRITE(OUTUNIT, 405) APtitle, imod, IPreCalcOutput,
                       RelDwellTime(Ilbins), XLbounds(ILbins),
      #
                       XLbounds (Ilbins+1)
     #
         READ(INPUNIT, 1411) Norbits, Norbsteps
         WRITE (OUTUNIT, 411) Norbits, Norbsteps
         READ(INPUNIT, 1412) OrbPeriodHrs
         WRITE(OUTUNIT,412) OrbPeriodHrs
         READ (INPUNIT, 9194) TITLELINE
         WRITE (OUTUNIT, 9194) TITLELINE
         READ (INPUNIT, 9194) TITLELINE
         WRITE (OUTUNIT, 9194) TITLELINE
         NE = 29 !presently hardwired for pre-calculated fluxes.
         DO I=1, NE, 2
             READ(INPUNIT, 9201) Evals(I), PCfluxpts(I),
             DiffPCfluxpts(I), Evals(I+1), PCfluxpts(I+1),
      #
             DiffPCfluxpts(I+1)
             WRITE (OUTUNIT, 9200) (Evals (K), PCfluxpts (K),
             DiffPCfluxpts(K), K=I, I+1)
```

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```
READ(INPUNIT, 9100) ELOWER, EUPPER, MPTS, IZLOW, IZHIGH, TARGET,
                 YEAR, NE, VERSION TMP, CODE TMP
          READ (INPUNIT, 100)
          WRITE(OUTUNIT, 9100) ELOWER, EUPPER, MPTS, IZLOW, IZHIGH, TARGET,
                 YEAR, NE, VERSION NUMBER, PROGRAM CODE
          WRITE (OUTUNIT, 100)
          Write trapped proton fluxes to file in standard CREME96 format.
  C
          READ(INPUNIT, 200) (PCFluxes(K), K=1, Mpts)
          WRITE(OUTUNIT, 200) (PCFluxes(K), K=1, Mpts)
          READ (INPUNIT, 100)
          WRITE (OUTUNIT, 100)
          CLOSE (INPUNIT)
           CLOSE (OUTUNIT)
        ENDIF !for regular vs. pre-calculated trapped fluxes, 11-26-97.
I C
         FORMAT statements
        FORMAT (1X, 2 (1PE10.4, 2X), 3 (I5, 2X), A12,
  100
                2X, OPF8.3, 1x, I2, 1x, I1, 1x, I4, 1x, I1)
  200
       FORMAT((1X,6(1PE10.4,2X)))
   404 FORMAT(1x,'%Incl = ',F7.3,' deg Apo = ',E10.4,
                   ' Peri = ',E10.4,' km',1x,3(F6.2,1x))
       #
   405 FORMAT(1x,'%',A6,1X,'IMOD =',I2,1X,'IPRECALC =',I2,/,1X,
                   '%Relative dwell time = ',E10.4,1X,
                   'L Bin: ',2(E10.4,1X))
         Number of steps per orbits is presently fixed at 200, and
         the orbital procession period is not presently calculated, 11-26-97.
  С
    411 FORMAT(1x,'%No. orbits =', I8, 2X,'No. steps/orbit = ', I6)
  cf? 411 FORMAT(1x,'%No. orbits =', I8, 2X,'No. steps/orbit = ', I6,
                      'Precession Period = ',E10.4)
  cf?
    412 FORMAT(1x,'%Orbital period = ',F8.2,1X,'hours')
    990 FORMAT(I3,1x,A70,I4,1x,I1)
    992 FORMAT(1x,'%Created by TRAPPED_DRIVER Version', I4,
                      ' on ',A9,' at ',A8)
   9100 FORMAT (1X, 2 (1PE10.4, 2X), 3 (15, 2X), A12, 2X, 0PF8.3, 1X, I4, 1X, I4, 1X, I1)
   9195 FORMAT(1X,'%Calculated energies, integral fluxes, and ',
                   'differential fluxes')
   9196 FORMAT(1X,'%',3X,'MeV',7X,'/m**2-sr-s',2X,'/m**2-sr-s-MeV',3X,
                   'MeV',7X,'/m**2-sr-s',2X,'/m**2-sr-s-MeV')
   9200 FORMAT((1X,'%',1X,3(1PE10.4,2X),3X,3(1PE10.4,2X)))
   9210 FORMAT((1X,3(1PE10.4,2X)))
```

```
1404 FORMAT(1x,8X,F7.3,12X,E10.4,8X,E10.4,3X,1x,3(F6.2,1x))
   1405 FORMAT (1x, 1X, A6, 1X, 6X, I2, 1X, 10X, I2)
   1406 FORMAT (1X, 23X, E10.4, 1X, 7X, 2 (E10.4, 1X))
   1411 FORMAT(1x, 13X, I8, 2X, 18X, I6)
   1412 FORMAT (1x, 18X, F8.2, 1X, 5X)
   9194 FORMAT(1X,A79)
   9201 FORMAT((1X,1X,1X,3(1PE10.4,2X),3X,3(1PE10.4,2X)))
         RETURN
         END
           SUBROUTINE Trapped_ORBINT(OrbIncl,Apogee,Perigee,AscNodeLong,
                    AscNodeDisp, PerigDisp, Evals, NE, sumexp, d,
        #
                    Year, XLbounds, ILbins, imod, RelDwellTime,
        #
                    DistBelt, PreCalcFlux, IPreCalc, Ndays,
        #
                    Norbits, Norbsteps, Period, OrbPrecTime)
IMPLICIT NONE
           INTEGER J, Jmax, L, Ndays, NorbSteps, IPreCalc, NLvals, Norbits
           INTEGER NE, imod
           Establish Norbits & Norbsteps in TrappedDriverInput, 11-26-97.
           PARAMETER (NLvals=10)
LOGICAL DistBelt, PreCalcFlux
$<del>*</del>
Initial Orbital parameters are set in Subroutine TrappedDriverInput.
Li
           REAL OrbIncl, Apogee, Perigee, AscNodeLong, AscNodeDisp, PerigDisp
REAL Time, Period, Step
Ų.
            Parameters along each orbital step
            REAL Zlat, Zlon, Alt
            INTEGER ILbins, ILbin, ICODE, NperLbin (NLvals)
            REAL Year, XLval, BB0, XLbounds (NLvals), XLinfinite
            PARAMETER (XLinfinite=1.0E+06)
            INTEGER IE, iemax
            parameter (iemax = 30) ! MAX NO. OF ENERGIES ALLOWED
            real Evals(iemax)
            REAL RelDwellTime (NLvals)
            REAL flux(iemax), expose(iemax)
            REAL sumexp(iemax, NLvals), fluxi(iemax, NLvals)
            REAL difed1(iemax, NLvals), d(iemax, NLvals)
            real*4 b, delt !B is mag. field, delt=step (time interval)
```

real*4 yearp

real*8 gmagmo

FORMAT lines for reading pre-calculated files, 11-26-97 and 12-1-97.

```
common /gmagmo/ gmagmo! for esa traraln
        SAVE
        real OrbPrecTime !for future use, 12-1-97.
  С
          Initializations
          DO L=1, NL vals
           NperLbin(L)=0
            do ie=1,ne ! initailize arrays
               sumexp(ie,L) = 0.
               fluxi(ie,L) = 0.
            enddo
           ENDDO
            do ie=1,ne ! initailize arrays
               expose(ie) = 0.0
            enddo
           IF (PreCalcFlux) THEN
               The output routine OutputTrappedFlux handles getting the
₫ C
               pre-calculated trapped fluxes. We simply return with
С
               the proper flag.
             RETURN !could just use subsequent RETURN, since this IF statement
                     !skips all lines before the subsequent RETURN
ELSE !calculate trapped flux if not using pre-calculated ones
lui.
             Initialize Orbit routine
CALL Orbit (1, Period, ZLon, ZLat, Alt, Apogee, Perigee, OrbIncl,
.
Lasis
                          AscNodeLong, AscNodeDisp, PerigDisp)
tage C
             Compute the total number of steps in "Ndays" days if we make
             "Norbsteps" steps per orbit. Use 2 days and 200 steps per orbit
   C
   С
             presently.
             NDAYS=INT(Norbits*PERIOD/86400. + 1.0)
             JMAX=INT(Ndays* NorbSteps*86400./PERIOD + 1.5)
   C
   С
             Compute the step size in seconds.
   С
             STEP=PERIOD/FLOAT (NorbSteps)
             delt=STEP
   C
            DO J=1, JMAX
             time=FLOAT(j-1)*step
             CALL Orbit(2, Time, ZLon, ZLat, Alt, Apogee, Perigee, OrbIncl,
```

#

```
call blccoords(Zlat, Zlon, Alt, Year, imod, BBO, XLval, yearp, B)
          IF ( XLbounds(2) .LT. XLinfinite .OR.
               ( XLbounds(2) .GE. XLinfinite .AND.
     &
                 XLbounds(1) .GT. 0.0 ) THEN
            IF (XLval .GT. 99999.0) XLval=99999.0
            CALL GetLbin(XLval, XLbounds, ILbins, ILbin)
            IF (ILbin .GE. 1 .AND. ILbin .LE. NLvals)
                NperLbin(ILbin) = NperLbin(ILbin) +1
          ELSE
            If no L-bins are specified or 1 L-bin is specified
C
            and the lower bound is L = 0, use only the first
C
            element of the array. In this case, the following
C
            sum should equal JMAX once the stepping through the
С
            orbit is completed.
            ILbin=1
            NperLbin(Ilbin) = NperLbin(Ilbin) +1
          ENDIF
         Now pass B, L (XLval), imod, e , ne to subroutine trapped_protons,
         return integral flux and yearp
С
         call trapped_protons(B, XLval, yearp, imod, evals(1), flux(1), ne)
          IF (ILbin .GE. 1 .AND. ILbin .LE. NLvals) THEN
           do ie = 1, ne
              expose(ie) = flux(ie) * delt
              sumexp(ie,ilbin) = sumexp(ie,ilbin) + expose(ie)
           enddo
          ENDIF ! for within allowed Lbins
                   !for number of orbital steps (up to JMAX)
         ENDDO
         DO L=1, ILbins
           do ie = 1, ne
              sumexp(ie,L) = sumexp(ie,L)/(FLOAT(NperLbin(L))*delt)
           enddo
           RelDwellTime(L) =
                                FLOAT (NperLbin(L))/
                                    FLOAT (Jmax)
     æ
           do ie = 1, ne-1
              difed1(ie,L) = sumexp(ie,L) - sumexp(ie+1,L)
           enddo
           difed1(ne,L) = sumexp(ne,L)
         ENDDO !stepping through L-bins
         call differ(ne, evals, sumexp, difed1, d, Ilbins)
```

```
do L = 1, ILbins
         do ie = 1, ne
           IF (d(ie,L) . LE. 1.0E-20) d(ie,L) = 0.0
       enddo
             !for using either pre-calculated or directly-calculated Fluxes
      ENDIF
      RETURN
      END
C-----
      The following GetLbin routine is identical to the routine in GEOMAG96.
      In order to avoid linking with all of GEOMAG96, it is included here
С
      for now. Before releasing, this should be in a standalone module
С
      that can be called from either the GTRANS DRIVER or the TRAPPED DRIVER.
C
C-----
      SUBROUTINE GetLbin(XLval, XLbounds, ILbins, ILbin)
      IMPLICIT NONE
      INTEGER ILbins, ILbin, NLvals, L
      PARAMETER (NLvals=10)
      REAL XLval, XLbounds (NLvals)
      LOGICAL FindLbin
      No attempt is made to eliminate "unphysical" or "approximate"
      L-values using the ICODE returned from GET BLCOORDS, since any
      analyses using L-values are likely to handle these locations
      "as is", i.e. with the calculated L-value.
C-----
      FindLbin=.TRUE.
      ILbin=0
      DO L=1, ILbins
       IF (FindLbin) THEN
         IF (L .LT. NLvals) THEN
           IF ( (XLval .GE. XLbounds(L)) .AND.
           (XLval .LT. XLbounds(L+1)) ) THEN
    &
             ILbin=L
             FindLbin=, FALSE,
           ENDIF
         ELSE !special handling of L=NLvals case
           IF (XLval .GE. XLbounds(L)) THEN
             ILbin=L
             FindLbin=.FALSE.
           ENDIF
         ENDIF !checking of each L-bin
       ENDIF
                !for FINDLbin logical
       ENDDO
```

```
C-----
         This should be a dead subroutine now, 11-26-97.
         SUBROUTINE GetPreCalcFlux(IPreCalc, Evals, TrappedFlux,
                     DiffTrappedFlux, DistBelt, imod)
       &
         IMPLICIT NONE
         INTEGER IPreCalc, NLvals, NE, imod, I, L, iemax
         PARAMETER (iemax=30, NLvals=10)
         LOGICAL DistBelt
         Initial Orbital parameters are set in Subroutine TrappedDriverInput.
  C
         real Evals (iemax)
         REAL Trappedflux(iemax, NLvals), DiffTrappedflux(iemax, NLvals)
         REAL RelDwellTime (NLvals)
  C-----
         RelDwellTime(1)=1.0
         DO I=1, iemax
          DO L=2, NL vals
             Trappedflux(i,L)=0.0
             DiffTrappedflux(i,L)=0.0
           ENDDO
         ENDDO
         DO L=2, NL vals
           RelDwellTime (L) = 0.0
Li
         ENDDO
Need to set Evals, Trappedflux(*,1), & DiffTrappedflux(*,1)
  С
         when actually implement routine. Also change STOP TO RETURN
   С
         WRITE(*,'(1X,''Pre-calculated trapped fluxes are presently'',
          1X,''not available.'',/,1X,''Aborting Trapped_Driver'')')
         STOP !
                       RETURN
          RETURN
         END
   C-----
         SUBROUTINE Trapped Spectra (Evals, NE, ELOWER, EUPPER, M, ILbins,
                     TrappedFlux, DiffTrappedFlux, Eout, Fluxout)
          IMPLICIT NONE
```

INTEGER NLvals, IEmax, NE, ILbins
PARAMETER (NLvals=10, IEmax=30)

RETURN END

```
REAL Evals(IEmax), DiffTrappedFlux(IEmax, NLvals)
       REAL TrappedFlux(IEmax,NLvals)
       REAL DiffFluxSPin(IEmax)
       REAL ELOWER, EUPPER, DE
       INTEGER MARR, M, I, J, K, NEsp, K1
       PARAMETER (MARR=5000)
       REAL Eout (MARR), Fluxout (MARR, NLvals), DiffFluxSPout (MARR)
       REAL D2FLUX(500), DUMMYFLUX(500), DERIVLOW, DERIVHIGH
       LOGICAL NATURAL(2)
       INTEGER Nelogmin, Nelogmax, Nelog
       REAL D2LOGFLUX(500), DUMMYLOGFLUX(500)
       REAL DERIVLOGLOW, DERIVLOGHIGH
       REAL EvalsLog(IEmax), Elogout(MARR), DifffluxLogout(MARR)
       REAL DiffFluxlog(IEmax)
       INTEGER Jmax
                       !11-24-97
                               _____
    Compute energies on logaritmically-spaced grid
     M=1002
     DE=(EUPPER/ELOWER) ** (1./(M-1.))
     EOUT (1) = ELOWER
     ELOGOUT (1) = LOG (EOUT (1))
     DO J=2, M-1
       EOUT(J)=EOUT(J-1)*DE
     ELOGOUT(J) =LOG(EOUT(J))
     END DO
      EOUT (M) = EUPPER
      ELOGOUT (M) = LOG (EOUT (M) )
С
     Initialize FluxOut
      DO I = 1, ILbins
        DO J=1,M
          FluxOut(J,I)=0.0
        END DO
      END DO
      EOUT (M) = EUPPER
      NATURAL(1) = .TRUE.
      NATURAL(2) = .TRUE.
      DO I = 1, ILbins !set up and CALL SPLINE for each L-value bin.
        NEsp=0
        NElogmin=0
        NElogmax=0
        Jmax=0
        DO J=1,IEMAX !establish array for passing into SPLINE & SPLINT
```

```
DiffFluxSPin(J) =DiffTrappedFlux(J,I)
            Eliminate points at which differential flux is not well
С
            behaved (monotonically decreasing), 11-24-97.
С
          IF (DiffFluxSPin(J) .GT. 0.0) THEN
            Eliminate points at which differential flux=0, 11-24-97.
С
            Jmax=J
            NEsp=NEsp+1
            IF (NElogmin .EQ. 0) NElogmin=J
            DiffFluxLog(NEsp-Nelogmin+1) = LOG(DiffFluxSPin(J))
            EvalsLog(NEsp-Nelogmin+1) = LOG(Evals(J)) !J to NEsp, 11-24-97.
          ENDIF
        ENDDO
        NElogmax=NEsp
        NElog=NElogmax-NElogmin+1
        IF (Nesp .GT. 1) THEN
          CALL SPLINE (EvalsLog, DiffFluxLog, NElog, 500, NATURAL
                       ,0.0,0.0,D2LOGFLUX,DUMMYLOGFLUX)
                         !for calculating at the standard CREME96 energies.
          DO K=1, M
            CALL SPLINT(Evalslog, DiffFluxLog, D2LOGFLUX,
                         NElog,Elogout(K),DiffFluxLogout(K))
             IF (Eout(K) .LT. 1.0 .OR.
                 Eout(K) .GT. Evals(Jmax)) THEN !NEsp to J, 11-24-97.
     &
               Fluxout (K, I) = 0.0
             ELSE
               Fluxout(K,I) = EXP(DiffFluxlogout(K))
             ENDIF
                         !K=1,M
           ENDDO
        ENDIF
               !I=1,ILbins
       ENDDO
       RETURN
       END
```

```
ELOWER, EUPPER, M, IZLO, IZUP,
                           FLUX, LDUM,
        &
                           SPECT)
        &
   C Creates an LET spectrum in standard format from an energy
   C spectrum in standard format.
   C Inputs:
   C SLOWER, SUPPER = min, max LET values (in MeV-cm2/g -- not /mg!)
              = target material (CHARACTER*7, generally SILICON)
            Contains the energy spectra of up to NELM elements specified at
   C FLUX
             up to MARR energies
   C
   C ELOWER, EUPPER = min, max energy of particle spectra
                 = number of bins in energy spectra
   C M
                  = min, max atomic numbers in LET spectrum.
   C IZLO, IZUP
                  = number of bins in integral LET spectrum (</= LARR=1002)
   C LDUM
   C Outputs:
                  = output LET spectrum
   C SPECT
   IMPLICIT NONE
         INTEGER*4 MARR, NELM, LARR, M, IZLO, IZUP, LDUM, L, J, K, I, IK
         PARAMETER (MARR=5000, NELM=92, LARR=1002)
         REAL*4 FLUX (NELM, MARR), E (MARR), SP (NELM, MARR)
         REAL*4 SPECT(LARR), SL(LARR)
         REAL*4 AMASS
         COMMON/MASS/AMASS(109)
         CHARACTER TARGET*12
         REAL*4 SLOWER, SUPPER, ELOWER, EUPPER, DE, DS, RSP, FUN, XK, ADD
   С
        Construct list of energies
ž
         DE = (EUPPER/ELOWER) ** (1./(M-1.))
11
         E(1)=ELOWER
DO J=2, M-1
E(J) = E(J-1) *DE
l-i
         END DO
ull.
         E(M) = EUPPER
         L=LDUM
         IF (L.GT.LARR) L=LARR
         Construct list of stopping powers
    С
         DS=(SUPPER/SLOWER)**(1./(L-1.))
         SL(1)=SLOWER
         DO J=2,L
           SL(J) = SL(J-1) *DS
         END DO
         SL(L)=SUPPER
    C
         Now get table of stopping powers
         CALL UNLOAD STABLE (ELOWER, EUPPER, M, IZLO, IZUP, TARGET, SP)
    С
         Initialize spectrum
         DO J=1,L
           SPECT(J) = 0.
```

END DO

SUBROUTINE ULET96 (SLOWER, SUPPER, TARGET,

С

```
For each energy find stopping power index and increment
all lower stopping powers
RSP=1./SLOWER
FUN=1./LOG(SUPPER/SLOWER)
DO J=IZLO, IZUP
  DO K=1, M
    XK=1.+(L-1.)*LOG(SP(J,K)*RSP)*FUN
    IK=INT(XK)
    IF (K.EQ.1) THEN
      ADD=FLUX(J,K)*(E(K+1)-E(K))*0.5
    ELSE IF (K.EQ.M) THEN
      ADD=FLUX(J,K)*(E(K)-E(K-1))*0.5
    ELSE
      ADD=FLUX(J,K)*(E(K+1)-E(K-1))*0.5
    ENDIF
    DO I=1, IK
      IF (I.LT.IK) THEN
        SPECT(I) = SPECT(I) + ADD
      ELSE IF (I.EQ.IK) THEN
        SPECT(I) = SPECT(I) + ADD*(XK-IK)
      ENDIF
    END DO
  END DO
END DO
RETURN
END
```

```
SUBROUTINE UNLOAD CREME96 FLUX(INFILE,
                                         ELOWER, EUPPER, M, IZLO, IZUP,
        &
                                         FLUX)
         IMPLICIT NONE
         INTEGER*4 MARR, NELM
         PARAMETER (MARR=5000, NELM=92)
         REAL*4 FLUX (NELM, MARR)
         REAL*4 ELOWER, EUPPER
         INTEGER*4 M, IZLO, IZUP, J, K, IVER, KVER, KPROG, NHEADER
         INTEGER*4 STAT, CREME96 OPEN
         CHARACTER*80 INFILE, ILINE
         CALL CHECK CREME96 VERSION (INFILE, IVER)
         OPEN(UNIT=25,STATUS='OLD',READONLY,FILE='USER:'//INFILE)
   С
         stat = creme96_open(infile,'user',25,'old')
         IF (IVER.GE.102) THEN
         READ(25,*) NHEADER
         DO J=1, NHEADER
            READ(25,110) ILINE
    110 FORMAT (A80)
         ENDDO
         ENDIF
         READ(25,*) ELOWER, EUPPER, M, IZLO, IZUP
         READ(25,*)
         DO 100 J=IZLO, IZUP
           READ(25,*) (FLUX(J,K),K=1,M)
           READ(25,*)
L C
            WRITE(6,999) J,(FLUX(J,K),K=1,6)
            WRITE(6,999) J,(FLUX(J,K),K=997,1002)
          FORMAT (1X, I3, 6E11.4)
   999
   100 CONTINUE
CLOSE (UNIT=25)
ĿĿ
         RETURN
         END
```

```
SUBROUTINE UNLOAD CTABLE (ELOWER, EUPPER, N, NSP, IZLO, IZUP, TARGET,
                                    CC, SPLOSS)
   С
         Subroutine used by UPROPC to unload cross-section tables.
   С
         IMPLICIT NONE
         INTEGER*4 MARR, NELM, MCS, STAT, CREME96 OPEN
         PARAMETER (MARR=5000, NELM=92, MCS=10)
         REAL*4 CC (NELM, NELM, MCS)
         REAL*4 SPLOSS (NELM, MCS)
         REAL*4 ELOWER, EUPPER, ELOWER$, EUPPER$
         INTEGER*4 N, NSP, NABS, IZLO, IZUP, N$, NSP$, IZLO$, IZUP$, I, J, K
         CHARACTER*12 TARGET, TARGET$
         CHARACTER*80 CTABLEFILE, SPTABLEFILE
         DATA ELOWER$, EUPPER$, N$, IZLO$, IZUP$,
                TARGET$/0.,0.,0,0,0,' '/
         FORMAT Statements
   C
         FORMAT(1X,2(1PE10.4,2X),4(15,2X),A12,2X,1PE10.4)
   100
         NABS=ABS(N)
   С
         First, check standard table:
          CTABLEFILE='CREME96:CTABLE.STD'
   C
          SPTABLEFILE='CREME96:SPTABLE.STD'
   С
T.
          IF (IZLO.GT.28 .or. IZUP.GT.28) THEN
   С
               CTABLEFILE='CREME96:CTABLE.XTD'
               SPTABLEFILE='CREME96:SPTABLE.XTD'
   C
          ENDIF
inches.
          CTABLEFILE='CTABLE.STD'
æ
          SPTABLEFILE='SPTABLE.STD'
المعا
          IF (IZLO.GT.28 .or. IZUP.GT.28) THEN
T.
              CTABLEFILE='CTABLE.XTD'
              SPTABLEFILE='SPTABLE.XTD'
          ENDIF
OPEN(UNIT=36,STATUS='OLD',FILE=CTABLEFILE,READONLY,SHARED)
          stat = creme96 open(ctablefile,'cr96tables',36,'old')
          READ(36,100) ELOWER$, EUPPER$, N$, IZLO$, IZUP$, NSP$, TARGET$
            WRITE(6,100) ELOWER$, EUPPER$, N$, IZLO$, IZUP$, NSP$, TARGET$
   CC
          IF (ELOWER.EQ.ELOWER$ .AND. EUPPER.EQ.EUPPER$ .AND.
             NABS.EQ.N$ .AND. NSP.EQ.NSP$ .AND.
              TARGET.EQ.TARGET$ .AND.
         ۶
             (IZLO$.LE.IZLO .AND. IZLO.LE.IZUP$) .AND.
             (IZLO$.LE.IZUP .AND. IZUP.LE.IZUP$)) THEN
          Standard table contains the necessary information.
    C
           OPEN(UNIT=37,STATUS='OLD',FILE=SPTABLEFILE,READONLY,SHARED)
          stat = creme96_open(sptablefile,'cr96tables',37,'old')
          READ (37, 100)
            READ(37,100) ELOWER$, EUPPER$, N$, IZLO$, IZUP$, NSP$, TARGET$
    CC
            WRITE(6,100) ELOWER$, EUPPER$, N$, IZLO$, IZUP$, NSP$, TARGET$
    CC
           WRITE(6,999) CTABLEFILE(1:20), SPTABLEFILE(1:20), TARGET
     999 FORMAT ('Standard tables ', A20, 1x, A20,
               /,' of nuclear cross-sections',
         δ
                  ' in ',A12,' used for transport calculation.')
```

```
GOTO 500
      ENDIF
      Check if appropriate tables exist in USER area:
С
      CLOSE (36)
       OPEN(UNIT=36,STATUS='OLD',FILE='USER:CTABLE.DAT',READONLY,ERR=50)
С
      stat = creme96_open('ctable.dat','user',36,'old')
      if (stat .ne. 0) goto 50
      READ(36,100) ELOWER$, EUPPER$, N$, IZLO$, IZUP$, NSP$, TARGET$
      IF (ELOWER.EQ.ELOWER$ .AND. EUPPER.EQ.EUPPER$ .AND.
         NABS.EQ.N$ .AND. NSP.EQ.NSP$ .AND.
           TARGET.EQ.TARGET$ .AND.
          (IZLO$.LE.IZLO .AND. IZLO.LE.IZUP$) .AND.
          (IZLO$.LE.IZUP .AND. IZUP.LE.IZUP$)) THEN
      Standard table contains the necessary information.
C
       OPEN(UNIT=37,STATUS='OLD',FILE='USER:SPTABLE.DAT',READONLY,ERR=50)
       stat = creme96_open('sptable.dat','user',37,'old')
       if (stat .ne. 0) goto 50
       READ (37, 100)
       WRITE(6,998) TARGET
 998 FORMAT(' User tables (USER:CTABLE.DAT & SPTABLE.DAT) of'
              ' nuclear cross-sections'
            /,' in ',A12,' used for transport calculation.')
       GOTO 500
       ELSE
       CONTINUE
 50
C
       CLOSE (36)
       CLOSE (37)
       WRITE(6,997)
 997 FORMAT(' Non-standard energy or shielding'
               ' in transport calculation.',
             /,' Create new cross-section tables in USER area.')
       CALL CTABLE (ELOWER, EUPPER, NABS, NSP, IZLO, IZUP, TARGET)
       READ (36,100) EUPPER$, ELOWER$, N$, IZLO$, IZUP$, NSP$, TARGET$
        OPEN(UNIT=37,STATUS='OLD',FILE='USER:SPTABLE.DAT',READONLY)
C
       stat = creme96 open('sptable.dat','user',37,'old')
       READ(37,100)
       ENDIF
  500 CONTINUE
       READ (36,100)
       READ (37, 100)
         DO J=1, NABS
            \texttt{READ} (\texttt{36}, \texttt{*}) \quad ((\texttt{CC}(\texttt{K}, \texttt{I}, \texttt{J}), \texttt{K} = \texttt{IZLO}\$, \texttt{IZUP}\$), \texttt{I} = \texttt{IZLO}\$, \texttt{IZUP}\$)
            READ(37,*) (SPLOSS(K,J),K=IZLO$,IZUP$)
          END DO
       CLOSE (UNIT=36)
       CLOSE (UNIT=37)
       RETURN
       END
```

#

la Fil

```
SUBROUTINE UNLOAD_HEADERS(INFILE,NHMAX,HEADER_LINE,LINEMAX)
С
      Reads lines of header information from file INFILE and
С
     returns LINEMAX lines in array HEADER_LINE. Maximum number
С
      of returned lines set by input NHMAX.
      IMPLICIT NONE
      CHARACTER*80 INFILE, HEADER_LINE
      DIMENSION HEADER_LINE(1)
      INTEGER*4 NHMAX, NHEADERS, VERSION_NUMBER, LINEMAX
      INTEGER*4 STAT, CREME96 OPEN
      INTEGER*4 INUNIT,J
      DATA INUNIT/4/
      CALL CHECK_CREME96_VERSION(INFILE, VERSION_NUMBER)
      CALL CHECK HEADER LENGTH (INFILE, NHEADERS)
      OPEN (UNIT=INUNIT, FILE='USER:'//INFILE,
С
            STATUS='OLD', READONLY, SHARED)
      stat = creme96_open(infile,'user',inunit,'old')
      By pass first line:
C
      IF (VERSION_NUMBER.GE.102) READ(INUNIT,999)
      Now store headers:
      LINEMAX=MIN(NHMAX,NHEADERS)
      DO J=1,LINEMAX
      READ(INUNIT, 999) HEADER_LINE(J)
 999 FORMAT (A80)
      ENDDO
      CLOSE (INUNIT)
      RETURN
      END
```

```
С
          Subroutine to unload integral LET Spectrum from input file into
  C
          array. Can handle either CREME96 format or the old CREME format
  С
          (ie., two-column table of LET (in MeV-cm2/g) and particle fluxes
  C
          (in particles/m2/s/sr).) CREME96 format is denoted by the
  C
          suffix .LET or .DLT in the filename.
  С
  С
                         Allan J. Tylka
          Written by:
  С
                         Code 7654
  С
                         Naval Research Laboratory
  C
                         Washington, DC 20375-5352
  C
                         tylka@crs2.nrl.navy.mil
  С
  С
          Last update: 31 October 1996: modified to read differential LET files.
  С
  С
  C-----
           IMPLICIT NONE
           INTEGER*4 K,N,NZ,NZT,I,NPTS,ILONG,MAXSPEC,STAT,CREME96 OPEN
           INTEGER*4 IVER, J, NHEADER
          PARAMETER (MAXSPEC=5000)
           CHARACTER*80 LETFILE, ILINE
          REAL*4 XL, FLUX, DUMFLUX, DUMXL, EL, EU
          DIMENSION XL(1), FLUX(1)
And the same
          DIMENSION DUMFLUX (MAXSPEC), DUMXL (MAXSPEC)
          DO 1 I=1, MAXSPEC
             DUMFLUX(I) = 0.0
          CONTINUE
   1
CALL CHECK_CREME96_VERSION(LETFILE, IVER)
<u>_</u>
₽.
           OPEN(UNIT=10,FILE='USER:'//LETFILE,STATUS='OLD',READONLY)
E C
           stat = creme96 open(letfile, 'user', 10, 'old')
ILONG=INDEX(LETFILE, '.')
Li
<u>l</u>
ų.
           IF (LETFILE(ILONG+1:ILONG+3).EQ.'LET' .or.
            LETFILE(ILONG+1:ILONG+3).EQ.'let' .or.
             LETFILE(ILONG+1:ILONG+3).EQ.'DLT' .or.
              LETFILE(ILONG+1:ILONG+3).EQ.'dlt') THEN
        δz
               IF (IVER.GE.102) THEN
               READ(10,*) NHEADER
               DO J=1, NHEADER
                  READ(10,110) ILINE
    110
                  FORMAT (A80)
               ENDDO
               ENDIF
               read(10,*) el,eu,n,nz,nzt
               FORMAT((1X,6(1PE10.4,2X)))
   200
               Calculate abscissae (LET values)
               DUMXL(1)=e1
               DUMXL(N)=eu
               do 100 i=2, N-1
```

DUMXL(i) = el*(eu/el)**(float(i-1)/float(n-1))

SUBROUTINE UNLOAD_LET_SPECTRUM(LETFILE,XL,FLUX,NPTS)

```
continue
   100
                Read blank line
   С
                read(10,110) ILINE
                read in the integral LET spectrum
   С
                read(10,200) (dumflux(i), i=1,n)
                CLOSE(10)
           ELSE
                Two-column format (not CREME96 standard)
   С
                N=1
                CONTINUE
    10
                READ(10, *, END=15) DUMXL(N), DUMFLUX(N)
                N=N+1
                GOTO 10
                CONTINUE
    15
                N=N-1
                CLOSE(10)
            ENDIF
   С
            Editing of input LET spectrum removed by AJT 10-21-96.
   С
            NPTS=0
            DO 1000 K=1, N
                        NPTS=NPTS+1
                        XL(NPTS) = DUMXL(K)
                        FLUX (NPTS) = DUMFLUX (K)
    1000
            CONTINUE
            RETURN
.
James
Harry Harry
            end
```

```
ELOWER, EUPPER, M, IZLO, IZUP,
        &
                                          FLUX)
   C
         From specified file INFILE unloads only elements IZMIN le Z le IZMAX
   С
         and returns the spectra in array FLUX.
   С
   С
         IMPLICIT NONE
         INTEGER*4 MARR, NELM, STAT, CREME96 OPEN
         PARAMETER (MARR=5000, NELM=92)
         REAL*4 FLUX(NELM, MARR), FLUXDUM(NELM, MARR), E(MARR)
         REAL*4 ELOWER, EUPPER, EMINCUT, EMAXCUT, DE
         INTEGER*4 IZMIN, IZMAX, M, IZLO, IZUP, KZLO, KZUP, J, K
         INTEGER*4 KMIN, KMAX, KMIN1, KMAX1
         INTEGER*4 IVER, NHEADER
         CHARACTER*80 INFILE, ILINE
         CALL CHECK_CREME96_VERSION(INFILE, IVER)
          OPEN(UNIT=25,STATUS='OLD',READONLY,FILE='USER:'//INFILE)
         stat = creme96_open(infile,'user',25,'old')
         IF (IVER.GE.102) THEN
         READ(25,*) NHEADER
         DO J=1, NHEADER
            READ(25,110) ILINE
   110 FORMAT(A80)
         ENDDO
Ц
         ENDIF
         READ(25,*) ELOWER, EUPPER, M, KZLO, KZUP
2
         READ(25,*)
<u>L</u>
         DO 100 J=KZLO,KZUP
FU.
          READ(25,*) (FLUXDUM(J,K),K=1,M)
READ(25,*)
    100 CONTINUE
         CLOSE (UNIT=25)
          IF (IZMIN.EQ.0) THEN
               IZLO=KZLO
          ELSE
               IZLO=MAX(IZMIN, KZLO)
          ENDIF
          IF (IZMAX.EQ.0) THEN
               IZUP=KZUP
          ELSE
               IZUP=MIN(IZMAX, KZUP)
          ENDIF
    С
          Now check energy limits:
          DE=(EUPPER/ELOWER) **(1./(M-1.))
```

IF (ELOWER.GE.EMINCUT) THEN

SUBROUTINE UNLOAD_PARTIAL_FLUX(INFILE, IZMIN, IZMAX, EMINCUT, EMAXCUT,

```
KMIN=1
    ELSE
        KMIN=1+IFIX(ALOG(EMINCUT/ELOWER)/ALOG(DE))
    ENDIF
    IF (EUPPER.LE.EMAXCUT) THEN
        KMAX=M
    ELSE
        KMAX=2+IFIX(ALOG(EMAXCUT/ELOWER)/ALOG(DE))
        IF (KMAX.GT.M) KMAX=M
    ENDIF
    KMIN1=MIN(KMIN,KMAX)
    KMAX1=MAX(KMIN,KMAX)
    DO 200 J=IZLO,IZUP
       DO 150 K=KMIN1,KMAX1
          FLUX(J,K) = FLUXDUM(J,K)
150
        CONTINUE
200 CONTINUE
```

RETURN END

```
SUBROUTINE UNLOAD PATH (IPATH, UPATH, TARGET, PATH, PSTEPMIN, PSTEPMAX,
                                PSTEP)
   С
         IMPLICIT NONE
         INTEGER*4 IPATH, NSTEP
         CHARACTER*12 TARGET
         REAL*4 UPATH, PATH, PSTEP, PSTEPMIN, PSTEPMAX, ALDEN
         DATA ALDEN/2.702000/
   C
        NOTE: current version of UNLOAD_PATH supports Al shielding only.
   С
   С
         IF (TARGET.NE.'ALUMINUM' .and. TARGET.NE.'aluminum') THEN
             WRITE (6, 9999)
             FORMAT(1X,' Specified shielding material unknown. STOP')
    9999
         ENDIF
         Convert input UPATH (which can be in mils, cm, or g/cm2 Al)
   C
         to g/cm2 Al.
   С
   С
         IF (IPATH.LT.0 .or. IPATH.GT.2) THEN
             WRITE(6,9998) IPATH
              FORMAT ('@ 04001 ABNORMAL TERMINATION: ',
    9998
               /,1x,' ERROR in UNLOAD PATH: ',
              /,1x,' PATH UNITS STEERING CODE UNKNOWN: ',15,
               /,1x,' STOP.')
             STOP
         ENDIF
         IF (IPATH.EQ.0) THEN
             Already specified in g/cm2
35
             PATH=UPATH
-
         ELSEIF (IPATH.EQ.2) THEN
             Specified in cm:
   C
             PATH=UPATH*ALDEN
ELSEIF (IPATH.EQ.1) THEN
             Specified in mils:
   С
             PATH=ALDEN*2.54*UPATH/1000.
         ENDIF
   C
         Now set PSTEP for transport.
            PSTEP=PSTEPMIN
            CONTINUE
    100
            NSTEP=PATH/PSTEP
            IF (NSTEP.GT.20) THEN
                PSTEP=PSTEP+0.10
                 GOTO 100
             IF (PSTEP.GT.PSTEPMAX) PSTEP=PSTEPMAX
   C
         Allow for very short PATHs:
          IF (PATH.LT.PSTEP) PSTEP=PATH
          RETURN
          END
```

```
С
          Subroutine to unload proton Spectrum from input file into
  C
          array. Can handle either CREME96 format or the old CREME format
  C
          (ie., two-column table of energies (in MeV) and fluxes (in
  C
          protons/m2-s-sr-MeV)). CREME96 format denoted .flx, .tfx, or
  C
  C
          .trp
  С
  С
                         Allan J. Tylka
  С
          Written by:
                         Code 7654
  C
  C
                         Naval Research Laboratory
                         Washington, DC 20375-5352
  C
                         tylka@crs2.nrl.navy.mil
  С
  C
          Last update: 17 November 1997: add .trp option
  С
  С
  С
       _____
  C-
  C
  С
          IMPLICIT NONE
          CHARACTER*80 PROTON FILE, ILINE
          INTEGER*4 MAXSPEC, N, NZ, NZT, MM, I, NPTS, ILONG, IZ
          INTEGER*4 IVER, J, NHEADER, STAT, CREME96 OPEN
REAL*4 EN, FLUX, EL, EU, DUMFLUX
          PARAMETER (MAXSPEC=5000)
          DIMENSION EN(1), FLUX(1), DUMFLUX (MAXSPEC)
          CHARACTER*3 SUFFIX
          CALL CHECK CREME96 VERSION (PROTON_FILE, IVER)
OPEN(UNIT=10,FILE='USER:'//PROTON_FILE,STATUS='OLD',READONLY)
E
           stat = creme96_open(proton_file,'user',10,'old')
12
           ILONG=INDEX(PROTON_FILE,'.')
SUFFIX=PROTON FILE (ILONG+1:ILONG+3)
          CALL CAPITALIZE STRING(SUFFIX, 3)
<u>ļ</u>.
ıI.
  C
          Filename check re-written AJT 12-9-97
          IF (SUFFIX .EQ. 'TFX' .or.
                          .EQ. 'FLX' .or.
             SUFFIX
        &
              SUFFIX(1:2) .EQ. 'TR') THEN
               IF (IVER.GE.102) THEN
               READ(10,*) NHEADER
              DO J=1, NHEADER
                 READ(10,110) ILINE
                  FORMAT (A80)
    110
               ENDDO
               ENDIF
               read(10,*) el,eu,n,nz,nzt
               IF (NZ.NE.1) THEN
               WRITE(6,999) PROTON FILE, NZ, NZT
    999
               FORMAT ('@ 09001 ABNORMAL TERMINATION: ',
                    /,1x,' ERROR IN UNLOAD_PROTON_SPECTRUM:',
                    /,1x,'
                           Specified file: ',
                    /,1x,A80,
                    /,1x,' includes ', I5,' .le. Z .le, ', I5,
        δ.
                    /,1x,' and contains no protons. STOP.')
```

SUBROUTINE UNLOAD PROTON SPECTRUM (PROTON_FILE, EN, FLUX, NPTS)

```
STOP
                                                                    ENDIF
            С
                                                                    Calculate abscissae (energy values)
            С
                                                                    EN(1)=el
                                                                    EN(N) = eu
                                                                    do 100 i=2, N-1
                                                                    EN(i) = el*(eu/el)**(float(i-1)/float(n-1))
                                                                    continue
             100
              С
                                                                    DO 150 IZ=NZ,NZT
                                                                    Read blank line
             C
                                                                    read(10,110) ILINE
                                                                    read in the flux
              С
                                                                    read(10,*)(dumflux(i),i=1,n)
                                                                    IF (IZ.EQ.1) THEN
                                                                                     DO 140 I=1,N
                                                                                                                       FLUX(I) = DUMFLUX(I)
 140
                                                                                      CONTINUE
the land don't have the corn of
                                                                    ENDIF
            150
                                                                    CONTINUE
                                                                    CLOSE (10)
                                                 ELSE
Mary offer from the state of th
                                                                    Standard old-CREME two-column format
                                                                    N=1
                  10
                                                                    CONTINUE
                                                                    READ(10, \star, END=15) EN(N), FLUX(N)
                                                                    N=N+1
                                                                    GOTO 10
                                                                    CONTINUE
                  15
                                                                    N=N-1
                                                                    CLOSE(10)
                                                  ENDIF
              С
                                                  Eliminate end-of-file zeroes from returned spectrum:
                                                  NPTS=0
                                                 DO 1000 I=1,N
                                                               IF (FLUX(I).GT.0.0) NPTS=I
                                                  CONTINUE
                  1000
                                                  RETURN
                                                  end
```

```
SUBROUTINE UNLOAD SHIELDFILE (SHIELDFILE,
                                          IUNITS, NSHIELD, UPATH, FRACSHLD)
         IMPLICIT NONE
         CHARACTER*80 SHIELDFILE
         INTEGER*4 MAXSHIELD, IUNITS, NSHIELD, K, STAT, CREME96 OPEN
         REAL*4 UPATH, FRACSHLD, TOTAL
         PARAMETER (MAXSHIELD=500)
         DIMENSION UPATH (MAXSHIELD), FRACSHLD (MAXSHIELD)
         INTEGER*4 ISHDUNIT/15/
         INTEGER*4 IVER, NHEADER
         CHARACTER*80 ILINE
         CALL CHECK CREME96 VERSION (SHIELDFILE, IVER)
         OPEN(UNIT=ISHDUNIT, READONLY, SHARED, STATUS='OLD',
   С
                FILE='USER:'//SHIELDFILE)
   С
         stat = creme96 open(shieldfile, 'user', ishdunit, 'old')
   C
         IF (IVER.GT.0) THEN
                READ (ISHDUNIT, *) NHEADER
                DO K=1, NHEADER
                   READ (ISHDUNIT, 5) ILINE
                   FORMAT (A80)
Ľ,
                ENDDO
The last than the last that
                READ (ISHDUNIT, *) IUNITS
         ENDIF
         K=0
         TOTAL=0
     10 CONTINUE
         K=K+1
IF (K.GT.MAXSHIELD) GOTO 100
         READ (ISHDUNIT, *, END=100) UPATH(K), FRACSHLD(K)
TOTAL=TOTAL+FRACSHLD(K)
13
         GOTO 10
14
    100 CONTINUE
<u>l</u>.,
         NSHIELD=K-1
         WRITE(6,999) NSHIELD
    999 FORMAT(' No. Shielding Bins = ', I4)
   C
         Renormalize shielding fraction to unit integral:
         DO 200 K=1, NSHIELD
                 {\tt FRACSHLD}\,({\tt K}) = {\tt FRACSHLD}\,({\tt K})\,/{\tt TOTAL}
    200 CONTINUE
         CLOSE (ISHDUNIT)
         RETURN
         END
```

```
SUBROUTINE UNLOAD STABLE (ELOWER, EUPPER, M, IZLO, IZUP, TARGET, SP)
         IMPLICIT NONE
         REAL*4 ELOWER, EUPPER
         INTEGER*4 M, IZLO, IZUP, NELM, MARR, STAT, CREME96 OPEN
         CHARACTER*12 TARGET
         PARAMETER (MARR=5000, NELM=92)
        REAL*4 SP(NELM, MARR)
        CHARACTER*12 TARGET$
        REAL*4 ELOWER$, EUPPER$
         INTEGER*4 M$, IZLO$, IZUP$
         INTEGER*4 J, K
         CHARACTER*80 STABLEFILE
        First, check standard table:
  С
        STABLEFILE='CREME96:STABLE.STD'
  C
         IF (IZLO.GT.28 .or. IZUP.GT.28) STABLEFILE='CREME96:STABLE.XTD'
  С
         STABLEFILE='STABLE.STD'
         IF (IZLO.GT.28 .or. IZUP.GT.28) STABLEFILE='STABLE.XTD'
         OPEN (UNIT=28, FILE=STABLEFILE, STATUS='OLD', READONLY, SHARED)
  С
         stat = creme96 open(stablefile,'cr96tables',28,'old')
        READ(28,100) ELOWER$, EUPPER$, M$, IZLO$, IZUP$, TARGET$
        IF (ELOWER.EQ.ELOWER$ .AND. EUPPER.EQ.EUPPER$ .AND. M.EQ.M$ .AND.
            TARGET.EQ.TARGET$ .AND.
        & (IZLO$.LE.IZLO .AND. IZLO.LE.IZUP$) .AND.
          (IZLO$.LE.IZUP .AND. IZUP.LE.IZUP$)) THEN
TI C
Standard table contains the necessary information.
  C
Ш
         WRITE(6,999) STABLEFILE(1:20), TARGET
<u>i</u>
   999 FORMAT(1x,' Standard table ', A20,' of stopping power',
               ' in ',A12,' used',/,1x,' for LET calculation.')
À
T.
         GOTO 500
Li
         ENDIF
<u>__</u>
  С
        Check if appropriate table exists in user area:
  C
        CLOSE (28)
         OPEN (UNIT=28, FILE='USER: STABLE.DAT', STATUS='OLD', READONLY, ERR=50)
   С
         stat = creme96_open('stable.dat','user',28,'old')
         if (stat .ne. 0) goto 50
         READ(28,100) ELOWER$, EUPPER$, M$, IZLO$, IZUP$, TARGET$
         IF (ELOWER.EQ.ELOWER$ .AND. EUPPER.EQ.EUPPER$ .AND. M.EQ.M$ .AND.
            TARGET.EQ.TARGET$ .AND.
            (IZLO$.LE.IZLO .AND. IZLO.LE.IZUP$) .AND.
            (IZLO$.LE.IZUP .AND. IZUP.LE.IZUP$)) THEN
   С
   C
        USER table contains the necessary information.
        WRITE(6,998) TARGET
    998 FORMAT(' User table (USER:STABLE.DAT) of stopping power'
            /,' in ',A12,' used for LET calculation.')
         GOTO 500
```

31

```
CLOSE (28)
                                          WRITE(6,997)
                   997 FORMAT(' Non-standard energy or material in LETSPEC calculation.',
                                                                 /,' Create new stopping power table in USER area.')
                                         CALL STABLE (ELOWER, EUPPER, M, IZLO, IZUP, TARGET)
                                              OPEN(UNIT=28,STATUS='OLD',FILE='USER:STABLE.DAT',READONLY)
                                          stat = creme96 open('stable.dat','user',28,'old')
                                          READ(28,100) ELOWER$, EUPPER$, M$, IZLO$, IZUP$, TARGET$
                                          ENDIF
                   500 CONTINUE
                                          READ (28, 100)
                                          DO J=IZLO$, IZUP$
                                                        READ (28,*) (SP(J,K),K=1,M$)
                                                        READ(28,*)
                                          ENDDO
                                           CLOSE (28)
The first that the first that
             100
                                          FORMAT(1X,2(1PE10.4,2X),3(I5,2X),A12,2X,1PE10.4)
                                          FORMAT((1X,6(1PE10.4,2X)))
             200
                                          RETURN
                                          END
 -
£.$
Specific Street, Specif
```

Create new STABLE.DAT in the users area:

ELSE CONTINUE

50 C

С

```
SUBROUTINE UNLOAD XSECT FILE (XSECT FILE, NSV, XV, YV)
С
С
        Subroutine to unload cross-section table from input file into
С
        array. Table is assumed to be in a two-column format, ordered
С
        according to increasing first-column value.
С
С
С
        Written by:
                       Allan J. Tylka
С
                       Code 7654
C
                       Naval Research Laboratory
С
                       Washington, DC 20375-5352
С
                       tylka@crs2.nrl.navy.mil
С
С
        Last update:
                       14 May 1996:
С
                       add 'USER:' to input file name.
С
C-
С
C
        IMPLICIT NONE
        CHARACTER*80 XSECT FILE
        INTEGER*4 NSV,I,STAT,CREME96_OPEN
        REAL*4 XV, YV
        DIMENSION XV(1), YV(1)
         OPEN(UNIT=10, FILE='USER:'//XSECT FILE, STATUS='OLD', READONLY)
        stat = creme96_open(xsect_file,'user',10,'old')
        I=1
        CONTINUE
10
        READ(10, *, END=15) XV(I), YV(I)
        I=I+1
        GOTO 10
        CONTINUE
 15
        NSV=I-1
        CLOSE(10)
        RETURN
        end
```

```
SUBROUTINE UNLOAD ZTABLE (ELOWER, EUPPER, M, IZLO, IZUP, TARGET, PSTEP,
  C
         Subroutine used by UPROPI to unload stopping power tables into array ZZ
  С
  C
         IMPLICIT NONE
         INTEGER*4 MARR, NELM, STAT, CREME96 OPEN
         PARAMETER (MARR=5000, NELM=92)
         REAL*4 ZZ (MARR, 2, NELM)
         REAL*4 ELOWER, EUPPER, PSTEP, ELOWER$, EUPPER$, PSTEP$
         REAL*4 DELTA PSTEP
         INTEGER*4 M, IZLO, IZUP, M$, IZLO$, IZUP$, MM, J, K
         CHARACTER*12 TARGET, TARGET$
         CHARACTER*80 ZTABLEFILE
  С
        FORMAT Statements
        FORMAT (1X, 2 (1PE10.4, 2X), 3 (15, 2X), A12, 2X, 1PE10.4)
  200
        FORMAT((1X,6(1PE10.4,2X)))
         First, check standard table:
  C
         ZTABLEFILE='CREME96: ZTABLE.STD'
  C
          IF (IZLO.GT.28 .or. IZUP.GT.28) ZTABLEFILE='CREME96:ZTABLE.XTD'
  С
         ZTABLEFILE='ZTABLE.STD'
...
         IF (IZLO.GT.28 .or. IZUP.GT.28) ZTABLEFILE='ZTABLE.XTD'
i c
         OPEN (UNIT=35, FILE=ZTABLEFILE, STATUS='OLD', READONLY, SHARED)
         stat = creme96 open(ztablefile,'cr96tables',35,'old')
         READ(35,100) ELOWER$, EUPPER$, M$, IZLO$, IZUP$, TARGET$, PSTEP$
        DELTA PSTEP=ABS (PSTEP-PSTEP$)
C
          TYPE *,' PSTEP, PSTEP$: ', PSTEP, PSTEP$
         IF (ELOWER.EQ.ELOWER$ .AND. EUPPER.EQ.EUPPER$ .AND. M.EQ.M$ .AND.
            TARGET.EQ.TARGET$ .AND. DELTA PSTEP.LE.0.001 .AND.
            (IZLO$.LE.IZLO .AND. IZLO.LE.IZUP$) .AND.
        & (IZLO$.LE.IZUP .AND. IZUP.LE.IZUP$)) THEN
IJ
         Standard table contains the necessary information.
Fill C
         WRITE(6,999) ZTABLEFILE(1:20), TARGET
    999 FORMAT(' Standard table ', A20,' of stopping power',
                ' in ',A12,' used for',/,' transport calculation.')
        &
         GOTO 500
         ENDIF
   C
   C
         Check if appropriate table exists in USER area:
         CLOSE (35)
         OPEN(UNIT=35, FILE='USER: ZTABLE.DAT', STATUS='OLD', READONLY, ERR=50)
   С
         stat = creme96 open('ztable.dat','user',35,'old')
         if (stat .ne. 0) goto 50
         READ (35,100) ELOWER$, EUPPER$, M$, IZLO$, IZUP$, TARGET$, PSTEP$
         DELTA PSTEP=ABS(PSTEP-PSTEP$)
         IF (ELOWER.EQ.ELOWER$ .AND. EUPPER.EQ.EUPPER$ .AND. M.EQ.M$ .AND.
             TARGET.EQ.TARGET$ .AND. DELTA PSTEP.LE.0.001 .AND.
        & (IZLO$.LE.IZLO .AND. IZLO.LE.IZUP$) .AND.
            (IZLO$.LE.IZUP .AND. IZUP.LE.IZUP$)) THEN
```

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```
User table contains the necessary information.
  С
        WRITE(6,998) TARGET
   998 FORMAT(' User table (USER: ZTABLE.DAT) of stopping power'
              /,' in ',A12,' used for transport calculation.')
        GOTO 500
        ELSE
        CONTINUE
   50
  С
        Create new ZTABLE.DAT in the users area:
  C
        CLOSE (35)
        WRITE(6,997)
   997 FORMAT(' Non-standard energy or shielding or PSTEP'
              ' in transport calculation.',
              /,' Create new stopping power table in USER area.')
        CALL ZTABLE (ELOWER, EUPPER, M, IZLO, IZUP, TARGET, PSTEP)
         OPEN(UNIT=35,STATUS='OLD',FILE='USER:ZTABLE.DAT',READONLY)
  С
        stat = creme96_open('ztable.dat','user',35,'old')
        READ(35,100) ELOWER$, EUPPER$, M$, IZLO$, IZUP$, TARGET$,
                 PSTEP$
ENDIF
   500 CONTINUE
        READ (35, 100)
        DO J=IZLO$, IZUP$
             READ(35,*) (ZZ(K,1,J),K=1,M$)
             READ(35,100)
ļad.
             READ (35, *) (ZZ(K, 2, J), K=1, M\$)
             READ(35,100)
END DO
T.
         CLOSE (UNIT=35)
ۇسۇ
ئ
         RETURN
         END
```

```
SUBROUTINE UPROP96 (INPUT FLUX,
                          ELOWER, EUPPER, M, IZLO, IZUP,
                          NDUM, NSP, PATH, PSTEP, TARGET,
       &
                          OUTPUT FLUX)
       &
  C This module evaluates nuclear transport, by calculating a numerical solution
  C to a one dimensional continuity equation, taking into account both
  C ionization energy loss (in the continuous-slowing-down approximation) and
  C nuclear fragmentation.
  C
  C This code is based on UPROP, originally written by John R. Letaw of Severn
  C Communications Corporation, working under contract to the Gamma Ray and
  C Cosmic Ray Astrophysics Branch of Naval Research Laboratory in 1989.
  C See "UPROP: A Heavy-Ion Propagation Code", by J.R. Letaw, SCC Report 89-02,
  C 31 August 1989.
  C
  C Adapted for use with CREME96 by AJT.
                                         Last Update 05 June 1996
  C Important Parameters
  C
          Maximum number of logarithically-spaced energy bins in spectrum
  C MARR
          Maximum atomic number of elements to be transported (<= 109)
C NELM
          Flag determining treatment of nuclear spallation reactions
              If NDUM=0 does not include nuclear spallation
  C
                  NDUM<0 does not follow nuclear fragments
三 C
                   |NDUM| = 1 uses energy-independent cross sections
TII C
                   |NDUM|>1 uses cross sections calculated at N equally-spaced
  С
                           energies and interpolated between
  C
  C PATH
           Total propagation pathlength in g/cm**2
  C PSTEP A small pathlength over which 2 nuclear fragmentations are
           unlikely, typically 0.1 g/cm**2.
  C TARGET Name of the target shielding material
  C INFILE File containing the heavy-ion energy spectrum (<= 40 bytes)
  C OUTFILE File containing the energy spectrum after transport (<= 40 bytes)
E C
C Important variables
  С
            working array contains the energy spectra of up to NELM elements
  C FLUX
            specified at up to MARR energies
  C
  IMPLICIT NONE
        INTEGER*4 MARR, NELM
        PARAMETER (MARR=5000, NELM=92)
        REAL*4 INPUT FLUX (NELM, MARR), OUTPUT FLUX (NELM, MARR)
        REAL*4 FLUX (NELM, MARR)
        REAL*4 ELOWER, EUPPER, PATH, PSTEP
        INTEGER*4 M, N, NDUM, IZLO, IZUP, J, K, NSP
        CHARACTER TARGET*12
  C
  C
        Copy input fluxes to working array:
        DO 20 J=IZLO, IZUP
           DO 10 K=1, M
              FLUX(J,K) = INPUT_FLUX(J,K)
   10
           CONTINUE
        CONTINUE
   20
```

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```
Special case for very thin shield (0.20 g/cm2 = 29.16 mils Al):
   С
   С
         Added by AJT, per JHA suggestion, 6-5-96
   С
         IF (PATH.LT.0.20) THEN
              CALL THIN_SHIELD(ELOWER, EUPPER, M, IZLO, IZUP, TARGET, PATH, FLUX)
         ELSE
         N=NDUM
         IF (PATH.LT.0.1) N=0
         IF (N.NE.O) THEN
         Alternate ionization loss and fragmentation using the pathlength
   С
         PSTEP until PATH is accumulated.
            DO J=1, INT (PATH/PSTEP+0.5)
              CALL UPROPI (ELOWER, EUPPER, M, IZLO, IZUP, TARGET, PSTEP, FLUX)
              CALL UPROPC (ELOWER, EUPPER, M, N, NSP, IZLO, IZUP, TARGET, PSTEP, FLUX)
            END DO
          ELSE
         Do ionization loss (only) using the pathlength PSTEP until PATH
   C
          is accumulated
for the first that the first first
            DO J=1, INT (PATH/PSTEP+0.5)
              CALL UPROPI (ELOWER, EUPPER, M, IZLO, IZUP, TARGET, PSTEP, FLUX)
            END DO
          ENDIF
į.i.
          ENDIF
ž
THE HOLD THE COL
          Copy transported energy spectra to output arrays:
          DO 200 J=IZLO, IZUP
             DO 100 K=1, M
                OUTPUT FLUX(J,K)=FLUX(J,K)
             CONTINUE
    100
    200
          CONTINUE
          RETURN
          END
```

```
PSTEP, FLUX)
  C SUBROUTINE UPROPC in Module UPROP.FOR
  C
  C Nuclear spallation subroutine. Determines the attenuation of a heavy-ion
  C energy spectrum from spallation reactions in passage through shielding
  C material. Initiates creation of an auxiliary data file (CTABLE.DAT) if
  C a suitable one does not already exist.
  C Modified by A.F. Barghouty 3-25-96
  C Parameters
           Maximum number of logarithically-spaced energy bins in spectrum
  C MARR
           Maximum atomic number of elements to be transported (<= 109)
  C NELM
           Maximum number of energies at which cross section data are
  C MCS
           defined
  C
  C ELOWER Lower energy bound of input and output spectra (>= 0.1 MeV)
  C EUPPER Upper energy bound of input and output spectra (<= 100000 MeV)
           Number of logarithmically equally-spaced energy bins (<= MARR)
  C M
           Number of energies at which cross section data are defined (<= MCS)
  C N
           Least atomic number of elements transported (>= 1)
  C IZLO
           Greatest atomic number of elements transported (<= 109)
  C IZUP
  C TARGET Name of the target shielding material (<= 12 bytes)
C PSTEP
           A small pathlength over which 2 nuclear fragmentations are
₫ C
           unlikely, typically 0.1 g/cm**2.
C FLUX
           Contains the energy spectra of elements IZLO through IZUP at
□ C
           M energies
TI C
  C Important variables
IJ C
           Energy at each M-point grid (spectrum grid)
  CE
           Energy at each N-point grid (cross section grid)
  C ECS
           Version number of current spallation cross section data file
  C VC
            (CTABLE, DAT)
  C
           Temporary flux vector used in calculating secondary spectra
  C FT
  C CC
           Partial and total cross section data computed by SPROP and stored
  С
           in CTABLE.DAT. First index is product Z; second index is target
A C
           Z; third index is index in N-point energy grid.
  C CCT
           Temporary array for holding cross sections at current M-point
  С
           energy grid.
  C REL
           Factor for normalizing energy to minimum energy on the grid
           Factor relating energy ratio to number of bins on the grid
  PARAMETER (MARR=5000, NELM=92, MCS=10)
        REAL*4 FLUX (NELM, MARR), CC (NELM, NELM, MCS), E (MARR), ECS (MCS)
        REAL*4 FT (NELM), CCT (NELM, NELM), SUM1 (NELM), SUM2 (NELM)
        REAL*4 SPLOSS (NELM, MCS), SPLT (NELM, MARR), dFLUX (NELM, MARR)
        REAL*4 FACTOR (NELM), TFLUX1 (NELM), TFLUX0 (NELM)
        CHARACTER*12 TARGET, TARGET$
        DATA IENT/0/
        IF (IENT.EQ.0) THEN
           IENT=1
           CALL UNLOAD CTABLE (ELOWER, EUPPER, N, NSP, IZLO, IZUP, TARGET,
                               CC, SPLOSS)
       &
        ENDIF
        NABS=ABS (N)
```

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SUBROUTINE UPROPC (ELOWER, EUPPER, M, N, NSP, IZLO, IZUP, TARGET,

```
С
         Compute energies associated with flux points
         REL=1./ELOWER
         FUL=1./LOG(EUPPER/ELOWER)
         DE=(EUPPER/ELOWER) ** (1./(FLOAT(M)-1.))
         E(1)=ELOWER
         DO I=2, M-1
           E(I)=E(I-1)*DE
         END DO
         E(M) = EUPPER
  С
         Compute energies associated with cross sections
         IF (NABS.GE.2) THEN
           DE=(EUPPER/ELOWER) ** (1./(FLOAT(NABS)-1.))
           ECS(1)=ELOWER
           DO I=2, NABS-1
             ECS(I) = ECS(I-1) *DE
           END DO
           ECS (NABS) = EUPPER
         END IF
  C
  С
         Perform fragmentation for each particle energy
□ C
1
        DO I=1, M
□ C
           Perform linear interpolation of cross
           section matrix appropriate for current energy
TI C
  С
ī c...
           [Interpolation debugged -for large II- 3/23/1996]
  С
IF (NABS.GE.2) THEN
200
             XI=1.+(NABS-1.)*LOG(E(I)*REL)*FUL
ļ.i.
             II=INT(XI)
T.
             IF (XI.GE.NABS) THEN
             DELTA=1./(ECS(NABS)-ECS(NABS-1))
             ELSE
               DELTA=1./(ECS(II+1)-ECS(II))
             ENDIF
             IF(I.LT.M) THEN
              F1X=DELTA*(E(I)-ECS(II))
              F2X=DELTA*(ECS(II+1)-E(I))
             END IF
             DO J=IZLO, IZUP
               DO K=IZLO, IZUP
                 IF(I.EO.1) THEN
                   CCT(J,K) = CC(J,K,1)
                    SPLT(J, I) = SPLOSS(J, 1)
                 END IF
                 IF (I.LT.M) THEN
                    CCT(J,K) = F1X*CC(J,K,II+1) + F2X*CC(J,K,II)
                    SPLT(J,I) = F1X*SPLOSS(J,II+1) + F2X*SPLOSS(J,II)
                 END IF
                 IF(I.EQ.M) THEN
                    CCT(J,K) = CC(J,K,NABS)
                    SPLT(J, I) = SPLOSS(J, NABS)
                 END IF
               END DO
```

END DO

```
DO J=IZLO, IZUP
               DO K=IZLO, IZUP
                 CCT(J,K) = CC(J,K,1)
                 SPLT(J,I) = SPLOSS(J,1)
               END DO
             END DO
           ENDIF
  C
           IF (N.GT.0) THEN
             If N > 0, compute fragmentation losses and gains
  C
             Form a temporary flux vector and multiply by PSTEP
  С
  C
             Secondaries (only) are computed from new vector
             DO J=IZLO, IZUP
               FT(J) = FLUX(J,I) * PSTEP
             END DO
  С
             Modify flux according to secondary production (includes
  С
  С
             all losses and gains)
             DO J=IZLO, IZUP
               DO K=IZLO, IZUP
                 FLUX(J,I) = FLUX(J,I) + CCT(J,K) * FT(K)
END DO
               IF (FLUX(J,I).LT.1.E-20) FLUX(J,I)=0.
             END DO
           ELSE IF (N.LT.0) THEN
             If N < 0, compute only fragmentation loss
#
DO J=IZLO, IZUP
FLUX(J,I) = FLUX(J,I) * (1.+CCT(J,J) * PSTEP)
W
               IF (FLUX(J,I).LT.1.E-20) FLUX(J,I)=0.
END DO
           ENDIF
         END DO
  С
  С
             Compute new flux taking into account energy losses due to
                                            (Sept. 1993)
   С
             fragmentation:
   С
             IF (NSP.EQ.1) THEN
              IF (IENT.EQ.O) THEN
              IENT=1
              WRITE (6,9999)
              FORMAT(1x,' UPROPC: Straight-ahead approx. NOT used.')
    9999
              END IF
   С
             DO I=IZLO, IZUP
                SUM1(I)=0.
                DO J=1,M
                  SUM1(I) = SUM1(I) + FLUX(I,J) *E(J)
                END DO
             END DO
   C
```

DO I=IZLO, IZUP

ELSE

```
DO J=1,M
                 dFLUX(I,J)=0.
   C
                 IF(J.EQ.1) THEN
                 dEN=1./(E(2)-E(1))
                 dFLUX(I,1) = dEN*(SPLT(I,1)*FLUX(I,2)+SPLT(I,2)*FLUX(I,1)
                             -2.*FLUX(I,1)*SPLT(I,1))
        &
   С
                 ELSE IF(J.EQ.M) THEN
                 dEN=1./(E(M)-E(M-1))
                 dFLUX(I,M) = dEN*(2.*FLUX(I,M)*SPLT(I,M)
        &
                             -SPLT(I, M-1) *FLUX(I, M) -SPLT(I, M) *FLUX(I, M-1))
   С
                 ELSE
                 dEN=1./(E(J+1)-E(J-1))
                 dFLUX(I,J) = dEN*((SPLT(I,J+1)-SPLT(I,J-1))*FLUX(I,J)
                                + (FLUX(I,J+1)-FLUX(I,J-1))*SPLT(I,J))
   С
                 END IF
   C
                 IF(ABS(dFLUX(I,J)).LT.1.E-20) dFLUX(I,J)=0.
                 FLUX(I,J) = FLUX(I,J) + dFLUX(I,J) * PSTEP
                 IF(FLUX(I,J).LT.1.E-20) FLUX(I,J)=0.
   С
END DO
               END DO
              DO I=IZLO, IZUP
                 SUM2(I)=0.
                 DO J=1,M
                  SUM2(I) = SUM2(I) + FLUX(I,J) *E(J)
H
              END DO
Œ
         .... Normalization:
řij.
             DO I=IZLO, IZUP
ZERO=ABS(1.-SUM1(I)/SUM2(I))
÷
                 IF (ZERO.GT..01) THEN
C C
                    TYPE *,' '
                    TYPE *,'
  С
                                *** Normalization...! ***'
   С
                    TYPE *,' '
                 END IF
              END DO
   C
              END IF
         RETURN
         END
```

```
C SUBROUTINE UPROPI in Module UPROP.FOR
  C Ionization loss subroutine. Determines the attenuation of a heavy-ion
  C energy spectrum from ionization losses in passage through shielding
  C material. Initiates creation of an auxiliary data file (ZTABLE.DAT) if
  C a suitable one does not already exist.
  C Parameters
           Maximum number of logarithically-spaced energy bins in spectrum
  C MARR
          Maximum atomic number of elements to be transported
  C NELM
  C ELOWER Lower energy bound of input and output spectra (<= 0.1 MeV)
  C EUPPER Upper energy bound of input and output spectra (>= 100000 MeV)
           Number of logarithmically equally-spaced energy bins (<= MARR)
  C M
  C IZLO
           Least atomic number of elements transported (>= 1)
  C IZUP
           Greatest atomic number of elements transported (<= 109)
  C TARGET Name of the target shielding material (<= 12 bytes)
  C PSTEP A small pathlength over which 2 nuclear fragmentations are
           unlikely, typically 0.1 g/cm**2.
  C FLUX
           Contains the energy spectra of elements IZLO through IZUP at
□ C
          M energies
₽ C
C Important variables
□ C
C FLUX2
           Temporary vector containing the energy spectrum of a single
  С
           element
U C ZZ
           Range and stopping power data computed by ZPROP and stored in
LL C
           ZTABLE.DAT
          Factor for normalizing energy to minimum energy on the grid
  C REL
           Factor relating energy ratio to number of bins on the grid
  C FUL
  IMPLICIT NONE
        INTEGER*4 MARR, NELM
        PARAMETER (MARR=5000, NELM=92)
        REAL*4 FLUX (NELM, MARR), ZZ (MARR, 2, NELM), E (MARR), FLUX2 (MARR)
        CHARACTER*12 TARGET
        INTEGER*4 IENT,M,IZLO,IZUP,I,J,K,KK
        REAL*4 ELOWER, EUPPER, PSTEP, REL, FUL, DE, XK
        DATA IENT/0/
        IF (IENT.EQ.0) THEN
           TENT=1
           CALL UNLOAD_ZTABLE (ELOWER, EUPPER, M, IZLO, IZUP, TARGET, PSTEP, ZZ)
        ENDIF
  C
        Compute new flux
        REL=1./ELOWER
        FUL=1./LOG(EUPPER/ELOWER)
        DE=(EUPPER/ELOWER)**(1,/(M-1.))
        E(1)=ELOWER
        DO I=2, M-1
          E(I)=E(I-1)*DE
        END DO
        E(M)=EUPPER
```

```
DO J=IZLO, IZUP
  DO K=1,M
     XK=1.+(M-1.)*LOG(ZZ(K,1,J)*REL)*FUL
     KK=INT(XK)
     IF (XK.GE.M) THEN
       FLUX2(K) = ((ZZ(K,1,J)-E(M-1))*FLUX(J,M)+
                (E(M)-ZZ(K,1,J))*FLUX(J,M-1))/(E(M)-E(M-1))
&
     ELSE
       FLUX2(K) = ((ZZ(K,1,J)-E(KK))*FLUX(J,KK+1)+
                (E(KK+1)-ZZ(K,1,J))*FLUX(J,KK))/(E(KK+1)-E(KK))
&
     ENDIF
     FLUX2(K) = FLUX2(K) * ZZ(K,2,J)
     IF (FLUX2(K).LT.1.E-20) FLUX2(K)=0.
   END DO
  DO K=1, M
     FLUX(J,K) = FLUX2(K)
   END DO
 END DO
 RETURN
 END
```

C

Module: VAX ROUTINES

С C С

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С

Logical Names and Environment Variables serve the same purpose, but are handled differently, on the two CREME96 platforms (VAX and PC respectively). There are also differences between the two file OPEN statements. To enable platform independance where fully specified filenames and where file opens are used in the higher level CREME96 code, two versions exist of the routines to perform these tasks. When an executable is being created, it is the responsibility of the person performing the link to ensure that the appropriate set of routines is used for the current build.

C С C С

Three platform-DEPENDANT routines exist:

C Ç

CREME96 FULL FILENAME CREME96 OPEN

creates fully-specified filename performs a file OPEN on full filename

C С С

С

CHECK FILE.FOR

Added 6/8/97

because VAX version contains LIB\$SPAWN

REMOVED 11/18/97 by AJT

С

SHOW DIRECTORY.FOR

Added 11/18/97; contains LIB\$SPAWN

These routines reside in the following 2 physical files:

VAX ROUTINES.FOR PC ROUTINES.FOR

used for a VAX build (this file)

used for a PC build

Integer function creme96 open(filename,path,unit,status)

C C

FILENAME:

The non-fully specified name of the target file.

С С

Contains the VMS logical pointing to directory PATH: where file does, or will exist.

С С С

UNIT:

The logical unit to be associated with the file. Must be defined at the time of the function call

(one will not be assinged by this routine).

С C С

С

STATUS:

Contains either OLD, for existing file, or

NEW, to create a file.

С С С

Calling example:

С С

STAT = creme96 open('input.dat','creme96',inunit,'old')

С С

Success is indicated by a ZERO return value. Otherwise, the return value will contain the FORTRAN error code.

С

IMPLICIT NONE

```
file, creme96 full filename, line
          character*80
                         filename
          character*(*)
          character*(*)
                         path
          integer
                         unit, ios
          character*(*)
                         status
          WRITE(*,*) 'IN OPEN... FILENAME: ',FILENAME,'
                                                       PATH: ',PATH
  С
          file = creme96 full filename(filename, path)
          if (status(1:1) .eq. 'o' .or. status(1:1) .eq. 'O') then
             Old files are only opened for READ (no APPEND in CREME96).
  С
             Any file opened for READ will be opened SHARED.
  С
             open (unit=unit, file=file, status='old',
             READONLY, SHARED, iostat=ios, err=199)
       &
  c DEBUG
             read(unit,99)line
  С
             format(a80)
  c99
             write(*,*)'First line in file: ',line
  С
          else
             New file to be created. WRITE and NOSHARE are default. On the PC,
□ c
             we must open with REPLACE instead of NEW, in case a file already
II c
             exists of this name (as it is our intention to write over it). If
E C
ij c
             one doesn't exist, REPLACE acts the same as NEW.
             OPEN (UNIT=unit, file=file, status='new',
             iostat=ios,err=199)
  c DEBUG
             write(*,*)'Writing test line to new file...'
  C
             write(unit,*)'Test line'
          endif
          creme96_open = ios
  199
          return
          end
   C
          Character*80 function creme96_full_filename(filename,path)
          The variable PATH contains the name of the VMS logical, which
   С
          in turn points to the directory path of the target file.
   С
          This routine appends the logical name to the bare filename.
   С
          IMPLICIT NONE
          character*(*)
                          filename
          character*(*)
                          path
          WRITE(*,*) 'IN FULL... FILENAME: ',FILENAME,'
                                                           PATH: ', PATH
          creme96 full filename = path//':'//filename
```

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```
WRITE(*,*) 'FULL FILENAME: ', CREME96 FULL FILENAME
          return
          end
  SUBROUTINE SHOW_DIRECTORY(JFILETYPE)
  C
  Ç
         VAX-specific routine.
  C
         INTEGER*4 JFILETYPE, ISTAT
         LOGICAL LIBSPAWN
         IF (JFILETYPE.EQ.0) THEN
          WRITE(6,9010)
          FORMAT(1x,' Here is the directory of your current USER area:')
   9010
          ISTAT=LIB$SPAWN('DIR USER:*.*')
         ELSEIF (JFILETYPE.EQ.1) THEN
          WRITE (6,9011)
          FORMAT(1x,' Here are the .TRP files in your current USER area:')
   9011
          ISTAT=LIB$SPAWN('DIR USER:*.TR*')
         ELSEIF (JFILETYPE.EQ.2) THEN
          WRITE(6,9012)
          FORMAT(1x,' Here are the .GTF files in your current USER area:')
   9012
ISTAT=LIB$SPAWN('DIR USER:*.GT*')
         ELSEIF (JFILETYPE.EQ.3) THEN
          WRITE(6,9013)
   9013 FORMAT(1x,' Here are the .FLX files in your current USER area:')
L
          ISTAT=LIB$SPAWN('DIR USER:*.FLX')
WRITE(6,9014)
          ISTAT=LIB$SPAWN('DIR USER:*.TFX')
WRITE(6,9011)
T.
          ISTAT=LIB$SPAWN('DIR USER:*.TR*')
L
ELSEIF (JFILETYPE.EQ.4) THEN
113
          WRITE(6,9014)
          FORMAT(1x,' Here are the .TFX files in your current USER area:')
   9014
          ISTAT=LIB$SPAWN('DIR USER:*.TFX')
          WRITE(6,9013)
          ISTAT=LIB$SPAWN('DIR USER:*.FLX')
          WRITE (6, 9011)
           ISTAT=LIB$SPAWN('DIR USER:*.TR*')
         ELSEIF (JFILETYPE.EQ.5) THEN
          WRITE (6,9015)
          FORMAT(1x,' Here are the .LET files in your current USER area:')
    9015
          ISTAT=LIB$SPAWN('DIR USER:*.LET')
         ELSEIF (JFILETYPE.EQ.6) THEN
          WRITE (6,9016)
          FORMAT(1x,' Here are the .DLT files in your current USER area:')
    9016
          ISTAT=LIB$SPAWN('DIR USER:*.DLT')
          ELSEIF (JFILETYPE.EQ.7) THEN
           WRITE (6, 9017)
          FORMAT(1x,' Here are the .SHD files in your current USER area:')
    9017
```

ISTAT=LIB\$SPAWN('DIR USER:*.SHD')

Œ

```
ELSEIF (JFILETYPE.EQ.8) THEN
WRITE(6,9018)

FORMAT(1x,' Here are the .XSD files in your current USER area:')
ISTAT=LIB$SPAWN('DIR USER:*.XSD')

ENDIF

WRITE(6,9999)

9999 FORMAT(/)
RETURN
END
```

```
REAL*4 FUNCTION WEIBULL (ONSET, WIDTH, POWER, ASYMPTOTE, E)
C
C
        Returns value of Weibull cross-section evaluated at abscissa E
C
        This function can be used for either heavy-ion or proton
C
C
        cross-sections; but the units are different in each case.
С
С
        Input parameters of Weibull fit:
C
                     (in MeV for proton; in MeV-cm2/mg for heavy ion LET)
          O= onset
С
           W= width
                       (as above)
С
           P= power
                       (dimensionless exponent)
C
          A= asymptote (in 10E-12 cm2/bit for protons cross-sections;
C
                        in square microns/bit for heavy-ion cross-sections)
          E= absicissa (in MeV for protons; in MeV-cm2/mg for heavy ion LET)
С
С
С
       Output: SEU cross-section (same units as asymptote)
С
C
       Written by:
                      Allan J. Tylka
C
                      Code 7654
С
                      Naval Research Laboratory
С
                      Washington, DC 20375-5352
С
                      tylka@crs2.nrl.navy.mil
С
С
       Last update:
                      29 March 1996
C
C-
   IMPLICIT NONE
       REAL*4 E, ONSET, Y, WIDTH, POWER, ASYMPTOTE
       WEIBULL=0
       IF (E.LT.ONSET) RETURN
       Y = ((E-ONSET)/WIDTH)**POWER
       Y=1.0-exp(-Y)
       WEIBULL=ASYMPTOTE*Y
       IF (WEIBULL.LT.O.) WEIBULL=0.
       RETURN
       END
```

```
C.....
   C... Silberberg&Tsao Semiempirical Cross Sections Routines.
   C... Notation:
   C... The S&T routines give the inelastic cross section QJ (in mb)
   C... for the reaction;
   C... (IZ,IA) + proton --> (JZ,JA) at energy EJ (in MeV/Nucleon),
   C... on the basis of a set of semiempirical formulae.
   C... Inputs: IZ - Atomic number of incoming nucleus
   C...
                IA - Atomic weight of incoming nucleus
                                                                . . .
   C...
                JZ - Atomic number of secondary nucleus
                JA - Atomic weight of secondary nucleus
   C...
   C...
                EJ - Amount of energy per nucleon of secondary nucleus...
   C... OUTPUT: QJ - Inelastic cross section of reaction
   C... References in commented lines are from:
   C... [1] Silberberg, R. & Tsao, C. H. 1973, Astrophys. J. Suppl.,
   C...
                                           25, pp. 315 - 333.
   C... [2] Silberberg, R. & Tsao, C. H. 1973, Astrophys. J. Suppl.,
   C. . .
                                           25, pp. 335 - 368.
   C... [3] Silberberg, R., Tsao, C. H. & Letaw, J. R.,
   C...
                       Astrophys. J. Suppl., 58, pp. 873 - 881.
   C...
  C... Another useful reference is:
  C...
           Silberberg, R. & Tsao, C. H. 1990, Phys. Rep., 191, 351.
  Ç...
   C...
                    *** Rewritten November 1995 ***
  C... Majority of comments written by Mark E. Mattson,
                                                                . . .
C... March - April, 1996, e-mail: mattson@vpihe4.phys.vt.edu
W C...
                                                                . . .
C... Routines linked:
  C... (1) PXN; for JZ = IZ+1 but JA=IA (i.e., pick-up reactions)
                                                                . . .
C... (2) YIELD1 (through 4 are spallation reactions); for JZ <= 4,
C... (3) YIELD2; for JZ >= 5 and 5 <= IZ <= 16
C... (4) YIELD3; for JZ >= 5 and 17 <= IZ <= 20
___ C... (5) YIELD4; for JZ >= 5 and 21 <= IZ <= 92
C... (6) CORRECTIONS; for various energy and/or structure-related
                                                                . . .
  C...
                       correction factors to spallation yields.
   C...
   C...........
   IMPLICIT NONE
        REAL*4 EJ,QJ
        INTEGER*4 IZ, IA, JZ, JA
        REAL*4 Q1,Q2,Q3,Q4,EXPF
        REAL*4 QR,QE,QF,QH,FE,FF,FA,FZ,CJ,PN,GA,ANZJ,AA,AE,AC,EC
```

LOGICAL NULL, NOT_CONSERVED, OUT_OF_RANGE, NON PICKUP, PICKUP, REJECTED LOGICAL REGION1, REGION2, REGION3

COMMON /FS/ QR,QE,QF,QH,FE,FF,FA,FZ,CJ,PN,GA,ANZJ,AA,AE,AC,EC

QJ = 0

C... Definitions:

C... This is in case the input indicates the secondary nucleus is exactly

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C... the same as the initial nucleus.
       NULL = IZ*IA.EQ.JZ*JA
 C... If the change in the atomic number of the nuclei is greater than the
 C... change in atomic number, then there's something wrong with the input.
       NOT_CONSERVED = ((IZ-JZ).GT.(IA-JA)) .OR. NULL
 C... If the incoming atomic weight is less than the secondary atomic weight,
 C... or if the incoming atomic number is greater than 92 (uranium), of if
 C... the incoming atomic number indicates hydrogen or helium, we're not
 C... interested in it.
      OUT_OF_RANGE = IA.LT.JA .OR. IZ.GT.92 .OR. IZ.LE.2
 C... If the secondary atomic number is 2 or more greater than the incoming
 C... atomic number, or if the number of neutrons in the secondary nucleus
C... is greater than the number of neutrons in the incoming nucleus, or if
C... the secondary nucleus is that of hydrogen, we're not interested in it.
      NON_PICKUP = (IZ-JZ).LT.-1 .OR. ((IA-IZ)-(JA-JZ)).LT.0
                                  .OR. JZ.LE.1
C... If the atomic number of the secondary is 1 greater than the atomic
C... number of the incoming and if the incoming atomic weight is at least
C... as great as the secondary atomic weight, we ARE interested in it, but
C... only as a (p,pxn) "pick-up" reaction.
      PICKUP = (IZ-JZ).EQ.-1.AND. IA.GE.JA
C... Regions of applicability as determined by the incoming nucleus:
      REGION1 = IZ.GE. 5 .AND. IZ.LE.16 ! Nuclei for Boron through Sulphur
      REGION2 = IZ.GE.17 .AND. IZ.LE.20 ! Nuclei for Chlorine through Calcium
      REGION3 = IZ.GE.21 .AND. IZ.LE.92 ! Nuclei for Scandium through Uranium
C... Initial Rejections:
      REJECTED = NULL.
     &
             OR.NOT CONSERVED.
              OR.OUT OF RANGE.
     &
     δe
              OR.NON PICKUP
      IF (REJECTED) RETURN
C... Non-Spallation, but pick-up reactions:
      IF (PICKUP) THEN
         CALL PXN(REAL(IZ), REAL(IA), REAL((IA-IZ)-(JA-JZ)), EJ,QJ)
         RETURN
      END IF
C... Spallation reactions sorted according to Z-number of secondary:
      IF (JZ.LE.4) THEN
                         ! If the secondary nucleus is between H and Be
         CALL YIELD1 (IZ, IA, JZ, JA, EJ, QJ)
      ELSE
         IF (REGION1) CALL YIELD2(IZ, IA, JZ, JA, EJ, QJ)
         IF (REGION2) CALL YIELD3(IZ, IA, JZ, JA, EJ, QJ)
         IF (REGION3) CALL YIELD4(IZ, IA, JZ, JA, EJ, QJ)
      END IF
C... Apply energy and/or structure-related correction factors:
      CALL CORRECTIONS (IZ, IA, JZ, JA, EJ, QJ)
     RETURN
     END
C
```

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C

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C
         SUBROUTINE CORRECTIONS (IZ, IA, JZ, JA, EJ, QJ)
    C.......
    C.....
   C. . .
   C... This subroutine includes corrections for both the energy and
   C... structure functions. Much of this is outlined in ref [3].
   C... Latest corrections, however, are outlined in Ref. 15 of CPC
   C... write-up!
   C...
   C.....
         IMPLICIT NONE
         REAL*4 EJ,QJ
         INTEGER*4 IZ, IA, JZ, JA
         REAL*4 Q1,Q2,Q3,Q4,EXPF
         REAL*4 QR,QE,QF,QH,FE,FF,FA,FZ,CJ,PN,GA,ANZJ,AA,AE,AC,EC
         COMMON /FS/ QR,QE,QF,QH,FE,FF,FA,FZ,CJ,PN,GA,ANZJ,AA,AE,AC,EC
   C... Energy-related corrections to results of YIELD1:
         IF (JZ.LE.4) THEN
           IF (IZ.EQ.6 .AND. EJ.GT.200.0 .AND. EJ.LE.400.0)
             QJ = QJ*(1.0 - 0.002*(EJ - 200.0))
           IF (IZ.EQ.6 .AND. EJ.GT.400.0 .AND. EJ.LE.1000.0)
             QJ = QJ*0.6
           IF (IZ.EQ.6 .AND. EJ.GT.1000.0 .AND. EJ.LE.5000.0)
             QJ = QJ*(0.6 + 0.0001*(EJ - 1000.0))
   C... Structure-related corrections to results of YIELD1:
           IF (IZ.EQ.6 .AND. IA.EQ.12 .AND. JZ.EQ.4 .AND. JA.EQ. 8)
             QJ = QJ*1.8
        ENDIF
C... Structure-related corrections to restuls of YIELD1, 3, and 4:
žesčis
        IF (IZ.EQ.7 .AND. IA.EQ.14 .AND. JZ.EQ.6 .AND. JA.EQ.12)
ű
             QJ = QJ*1.8
        IF (IZ.EQ.7 .AND. IA.EQ.14 .AND. JZ.EQ.6 .AND. JA.EQ.13)
             QJ = QJ*0.5
        IF (IZ.EQ.8 .AND. IA.EQ.16 .AND. JZ.EQ.6 .AND. JA.EQ.12)
             QJ = QJ*1.8
        IF (IZ.EQ.8 .AND. IA.EQ.16 .AND. JZ.EQ.7 .AND. JA.EQ.14)
             QJ = QJ*1.8
        IF (IZ.EQ.8 .AND. IA.EQ.16 .AND. JZ.EQ.7 .AND. JA.EQ.15)
             QJ = QJ*1.5
  С
  C...
  C
        IF (IZ.EQ. 8 .OR.IZ.EQ.10) THEN
           IF (IZ*2.EQ.IA.AND.JA-2*JZ.GE.2.AND.JZ.GE.5)
                                                      QJ=QJ*0.7
        ENDIF
        IF (IZ.GE. 9.AND.IZ.LE.16.AND.2*JZ-JA.EQ.1.AND.JZ.GE.9)QJ=QJ*.7
        IF (IZ.GE.10.AND.IZ.LE.13.AND.IA-IZ*2.NE.2)
            \label{eq:condition}  \text{IF((JZ.EQ.6.AND.JA.EQ.12).OR.(JZ.EQ.8.AND.JA.EQ.16))} \quad \text{QJ=QJ} \\ \times 2. 
        ENDIF
        IF((IZ.EQ.10.AND.IA.EQ.20).or.(IZ.EQ.12.AND.IA.EQ.24)) THEN
           IF (JZ.EQ.7.AND.(JA.EQ.14.OR.JA.EQ.15))
                                                           QJ=QJ*1.5
        ENDIF
```

C

C С C

C...

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IF (IZ.GE.10.AND.IZ.LE.16.AND.JZ.EQ.9 )
                                                         QJ=QJ*0.8
       IF ((IZ.EQ.12.OR.IZ.EQ.14.OR.IZ.EQ.16).AND.(2*IZ.EQ.IA)) THEN
          IF (IZ-JZ.EQ.2.AND.IA-JA.EQ.4)
                                                     QJ=QJ*1.6
         IF (IZ-JZ.EQ.1.AND.IA-JA.EQ.1)
                                                     QJ=QJ*1.6
      IF ((IZ.EQ.18.OR.IZ.EQ.20).AND.2*IZ.EQ.IA) THEN
        IF (IZ-JZ.EQ.1.OR.IZ-JZ.EQ.3)
                                                    QJ=QJ*0.7
      ENDIF
      IF (IZ.EQ.20.AND.IA.EQ.40.AND.(JZ.EQ.12.OR.JZ.EQ.14))QJ=OJ*2.4
      IF (IZ.EQ.20.AND.IA.EQ.40.AND.(JZ.EQ.18.OR.JZ.EQ.16))QJ=QJ*1.6
      IF (IZ.GE.24.AND.IZ.LE.28) THEN
         IF (JZ.GE.20.AND.JZ.LE.23.and.JA-JZ*2.GE.6)
                                                         QJ=QJ*0.5
      ENDIF
      IF((IZ.EQ.26.AND.IA.EQ.56).OR.(IZ.EQ.24.AND.IA.EQ.52)) THEN
         IF((JZ.EQ.20).OR.(JZ.EQ.18).OR.(JZ.EQ.16))
      ENDIF
      IF (IZ.GE.30.AND.IZ-JZ.GE.6)
     & QJ = QJ*(1.0+0.9*EXPF(-((EJ-1230)/150)**2)
                *EXPF(-(ABS(IZ-JZ-12)/5.)**2))
      IF (IZ.EQ.26 .AND. IA.EQ.56 .AND. JZ.EQ.23)
     & QJ = QJ*(1.0 - 0.6*EXPF(-((52-JA)/2.6)**2))
      IF (IZ.EQ.26 .AND. IA.EQ.56 .AND. JZ.EQ.24. AND. JA.EQ.54)
     & QJ = 0.7*QJ
      IF (IZ.EQ.26 .AND. IA.EQ.56 .AND. JZ.EQ.25. AND. JA.GE.54 .AND.
     & JA.LE.55) QJ = QJ*(1.7 - (JA-54)*0.45)
      IF (IZ.EQ.26 .AND. IA.EQ.56 .AND. JZ.EQ.26 .AND. JA.EQ.53)
     & QJ = QJ*2.0
      IF (IZ.EQ.26 .AND. IA.EQ.56 .AND. JZ.EQ.17) OJ = OJ*0.9
      IF (JZ.EQ.5) THEN
         CALL YIELD1 (IZ, IA, JZ, JA, EJ, Q1)
         QJ = SQRT(Q1*QJ)
         IF (IZ.EQ.7 .AND. IA.EQ.14 .AND. JZ.EQ.5 .AND. JA.EQ.10)
           QJ = QJ*1.8
         IF (IZ.EQ.6 .AND. IA.EQ.12 .AND. JZ.EQ.5 .AND. JA.EQ.10)
     æ
            QJ = QJ*1.8
         IF (IZ.EQ.6 .AND. IA.EQ.12 .AND. JZ.EQ.5 .AND. JA.EQ.11)
            QJ = QJ*1.5
     &
      ELSE
         IF (IZ.EQ.20.AND.JZ.GE.6) THEN
            CALL YIELD4 (IZ, IA, JZ, JA, EJ, Q4)
            QJ = SQRT(QJ*Q4)
         ENDIF
         IF (IZ.EQ.21 .AND. JZ.GE.6) THEN
            CALL YIELD3 (IZ, IA, JZ, JA, EJ, Q3)
            QJ = SQRT(QJ*Q3)
         ENDIF
      ENDIF
      RETURN
      END
C..........
C... The next four subroutines are used in the determination of the ...
C... various parameters in Table 1 of ref [1]. They are presented in...
C... this fashion to facilitate optimization with the data.
```

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```
C...
C...
             Y1BLOK - Parameters as in Table 1A of ref [1].
C...
             Y2BLOK - Parameters as in Table 1B of ref [1].
             Y3BLOK - Parameters as in Table 1C of ref [1].
C...
C...
             Y4BLOK - Parameters as in Table 1D of ref [1].
C...
C
      BLOCK DATA Y1BLOK
      COMMON /AT/ BA,C2,C3,C4,C5,QC,QL,B,PC,PL,RC,RL,RU,S,SE,C,T,ET
      DIMENSION BA(92), C2(56), C3(56), C4(56), C5(56), ET(4)
      DATA BA / 16 * 0.0, 36.0, 38.0, 40.0, 44.0,
     & 45.0, 48.0, 50.0, 52.0, 55.0, 55.7, 58.5, 61.0, 64.0, 67.0,
     & 70.0, 73.0, 75.0, 78.0, 80.0, 82.0,
                                             84.0, 86.0, 89.0, 92.0,
     & 93.0, 96.0, 98.0,100.0, 103.0, 106.0, 108.0, 111.0, 114.0, 118.0,
     &122.0,125.0,127.0,130.0, 133.0, 134.0, 137.0, 139.0, 141.0, 146.0,
     &146.0,149.0,153.0,156.0, 159.0, 161.0, 165.0, 166.0, 169.0, 172.0,
     &175.0,178.0,181.0,183.0, 186.0, 188.0, 192.0, 194.0, 197.0, 200.0,
     &204.0,206.0,209.0, 0.0, 0.0, 0.0, 226.0, 227.0, 232.0,
     &231.0,238.0/
C... Parameters for calculating OMEGA of Equation (1) in ref [1].
      DATA C2/5*1.0, 0.60, 1.00, 1.00, 1.00, 1.00, 46*1.0/
      DATA C3/5*1.0, 1.00, 1.80, 0.70, 3.00, 1.00, 46*1.0/
      DATA C4/6*1.0, 0.95, 1.00, 0.65, 1.30, 1.00, 45*1.0/
      DATA C5/ 7*1.0, 0.20, 1.00, 0.70, 1.70, 1.00, 44*1.0/
C... Parameters for calculating sigma 0 of Equation (1) in ref [1].
      DATA QC/ 13.0/, QL / 13.0/ , B / 1.15/
C... Parameters for calculating P of Equation (1) in ref [1].
      DATA PC/ 0.0/, PL / 0.16/
C... Parameters for calculating R of Equation (1) in ref [1].
      DATA RC/ 1.80/, RL / 10.7/ , RU/ 0.25/
C... Parameters for calculating S of Equation (1) in ref [1].
      DATA S / 0.54/, SE / 1.4/, C / 0.32/
C... Variable T of Equation (1) in ref [1].
      DATA T/ 0.003/
C... Parameter eta of Table 1A of ref [1].
      DATA ET / 1.15, 1.15, 0.9, 0.8/
      END
C
C
      BLOCK DATA Y2BLOK
      COMMON /BT/ QM, C5, C6, C7, C8, C9, CD, D1, D2, D3, D4, D5, D6, PC, PL, PU, PG, PH,
                     S, SE, C, RC, RL, RU, T, ET
C... The variable CJ holds the values for the parameter OMEGA of [1].
      DIMENSION ET(4), QM(7)
      DIMENSION C5(56), C6(56), C7(56), C8(56), C9(56), CD(56)
```

```
C... Parameters for determining sigma0 in Table 1B of ref [1].
        DATA QM / 27.6, 0.66667, 1.0, 1.0, 0.3, 0.05, 0.5/
   C... Parameters for determining P of [1].
        DATA PC, PL, PU/0.075, 2.60, 0.50/, PG, PH /0.77, 0.66667/
   C... Parameters for determining S of [1].
        DATA S, SE, C/0.502, 1.4, 0.26/
   C... Parameters for determining R of [1].
        DATA RC,RL,RU/1.60, 10.2, 0.26/
  C... The variable T in [1].
        DATA T/ 0.0005/
  C... Parameter eta as in Table 1B of ref [1].
        DATA ET / 1.15, 1.15, 0.9, 0.8/
ũ
END
  С
        BLOCK DATA Y3BLOK
COMMON /CT/ BA,QT,P0,P1,P2,P3,P4,P5,R0,R1,R2,S0,S1,T,ET
DIMENSION BA(92), QT(7), ET(4)
<u>...</u>
        DATA BA / 16 * 0.0, 36.0, 38.0, 40.0, 44.0,
H.
       & 45.0, 48.0, 50.0, 52.0, 55.0, 55.7, 58.5, 61.0, 64.0, 67.0,
& 70.0, 73.0, 75.0, 78.0, 80.0, 82.0, 84.0, 86.0, 89.0, 92.0,
Ŀ
       & 93.0, 96.0, 98.0,100.0, 103.0, 106.0, 108.0, 111.0, 114.0, 118.0,
&122.0,125.0,127.0,130.0, 133.0, 134.0, 137.0, 139.0, 141.0, 146.0,
*-
       &146.0,149.0,153.0,156.0, 159.0, 161.0, 165.0, 166.0, 169.0, 172.0,
       &175.0,178.0,181.0,183.0, 186.0, 188.0, 192.0, 194.0, 197.0, 200.0,
       &204.0,206.0,209.0, 0.0, 0.0, 0.0, 0.0, 226.0, 227.0, 232.0,
       &231.0,238.0/
  C... Parameters used in calculating sigma0 in Table 1C of ref [1].
        DATA QT/ 27.6, 0.6667, 1.0, 1.0, 0.3, 0.05, 0.5/
  C... Parameters used in calculating parameter P in Table 1C of ref [1].
        DATA P0,P1,P2,P3,P4,P5 /20.0, 0.77, 1.98, 0.92, 0.77, 0.6667/
  C... Parameters used in calculating parameter R in Table 1C of ref [1].
        DATA RO,R1,R2 /10.2, 0.26, 1.60/
  C... Parameters used in calculating parameter S in Table 1C of ref [1].
        DATA S0,S1 /0.502, 0.08/
```

DIMENSION D1(56), D2(56), D3(56), D4(56), D5(56), D6(56) DATA C5/ 7*1.0, 0.20, 1.00, 0.70, 1.70, 1.00, 44*1.0/

DATA C8/14*1.0, 1.20, 1.00, 1.00, 39*1.0/, C9/17*1.0,1.00,38*1.0/ DATA CD/18*1.0, 0.60, 1.00, 0.83, 35*1.0/, D1/21*1.0,1.20,34*1.0/

DATA D2,D3,D4,D5,D6/56*1.0, 56*1.0, 56*1.0, 56*1.0, 56*1.0/

DATA C6/ 9*1.0, 0.60, 46*1.0/

C... Parameters T in Table 1C of ref [1].

C... Values for eta in Table 1C of ref [1].

DATA T /0.0005/

DATA C7/12*1.0, 0.39, 1.00, 2.00, 41*1.0/

```
END
   C
   C
        BLOCK DATA Y4BLOK
        COMMON /DT/ BA,S0,S1,T3,R0,R1,R2,R3,P0,P1,P2,P3,E0,E1,C1,C2,C3,C4,
                   D1, D2, T2, ET
   C... BA is an array representing the average atomic number for all stable
   C... isotopes of a given element.
        DIMENSION
                  BA(92), ET(4)
        DATA BA / 16 * 0.0, 36.0, 38.0, 40.0, 44.0,
       & 45.0, 48.0, 50.0, 52.0, 55.0, 55.7, 58.5, 61.0, 64.0, 67.0,
       & 70.0, 73.0, 75.0, 78.0, 80.0, 82.0, 84.0, 86.0, 89.0, 92.0,
       & 93.0, 96.0, 98.0,100.0, 103.0, 106.0, 108.0, 111.0, 114.0, 118.0,
       &122.0,125.0,127.0,130.0, 133.0, 134.0, 137.0, 139.0, 141.0, 146.0,
       &146.0,149.0,153.0,156.0, 159.0, 161.0, 165.0, 166.0, 169.0, 172.0,
       &175.0,178.0,181.0,183.0, 186.0, 188.0, 192.0, 194.0, 197.0, 200.0,
       &204.0,206.0,209.0, 0.0, 0.0, 0.0, 226.0, 227.0, 232.0,
       &231.0,238./
   C... Parameters for calculating S, ?????? and R as in Table 1D of ref [1].
        DATA S0,S1,T3/0.482,0.07,3.0E-7/, R0,R1,R2,R3/11.8,0.45,1.29,0.15/
        DATA PO,P1,P2,P3/1.98, 0.92, 20.0, 0.77/, E0,E1/20.3, 1.169/
  C... Parameters for calculating sigma0 in Table 1D of ref [1] and Delta {\tt A\_c}
   C... in Equation (2) of ref [1].
        DATA C1,C2,C3,C4/144.0, 0.367, 0.3, 0.7/, D1,D2/0.0365, 1.23/
C... T in Table 1D of ref [1].
       DATA T2/ 2.8E-4/
C... Parameter eta in Table 1D of ref [1].
       DATA ET / 1.25, 0.9, 1.0, 0.85/
        END
  C
  C
  C
  C
       SUBROUTINE YIELD1 (IZ, IA, JZ, JA, EJ, QJ)
  C.....
  C.....
  C...
  C... This subroutine is for the case where the incoming nucleus is
  C... at least as large is lithium and the secondary nucleus is
  C... between helium and beryllium, i.e.,
  C...
            5 .LE. IZ .LE. 16
                             .AND. 5 .LE. JZ .LE IZ
  C...
  C.....
  C... Get parameters as in Table 1! of ref [1].
       EXTERNAL Y1BLOK
       COMMON /FS/ QR,QE,QF,QH,FE,FF,FA,FZ,PJ,PN,GA,ANZJ,AA,AE,AC,EC
```

DATA ET /1.25, 0.9, 1.0, 0.85/

COMMON /QG/ QI,G1,G2,G3,G4

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COMMON /ST/ ST, SS, T
      COMMON /AT/ BA,C2,C3,C4,C5,QC,QL,B,PC,PL,RC,RL,RU,S,SE,C,T,ET
      DIMENSION BA(92), CJ(56,26), C2(56), C3(56), C4(56), C5(56), ET(4)
C... Parameters for calculating OMEGA of Equation (1) in ref [1].
      EQUIVALENCE (CJ(1, 2), C2), (CJ(1, 3), C3), (CJ(1, 4), C4), (CJ(1, 5), C5)
      OR = 0.0
      QE = 0.0
      QF = 0.0
      QH = 0.0
      AE = 0.0
      AC = 0.0
      FE = 1.0
                ! Initial value of f(E) of Equation (1) of ref [1].
      FF = 1.0
      FA = 1.0
                ! Initial value of f(A) of Equation (2) of ref [1].
      FZ = 1.0
      PN = 1.0
      GA = 1.0
C... REAL*4 values for the integers describing atomic no. and atomic wqt.
      AI = IA
      ZI = IZ
      AJ = JA
      ZJ = JZ
C... Difference in atomic wgts.
      AA = AI - AJ
C... REAL*4 Number of neutrons in secondary nucleus.
      AN = AJ - ZJ
C... Ratio of neutrons to protons in secondary nucleus.
      ANZJ = AN/ZJ
C... Determination of the nuclear structure function, OMEGA, of Equation (1)
C... in ref [1].
      PJ = CJ(JA, JZ)
C... Determination of ratio (N/Z)* as on p. 349 of ref [2]; used in
C... calculation of f(A) of equation (1) of ref [1].
     \cdot AM = BA(IZ)
      IF (AM.EQ.0) AM = IA
      CN = 0.3*(AI - AM)/ZI
C... Change in the number of neutrons between target and secondary nuclei.
      KN = (IA - IZ) - (JA - JZ)
C... Change in the number of protons between target and secondary nuclei,
C... including initial proton.
      JP = IZ - JZ + 1
C... Integer number of neutrons in secondary nucleus.
      JN = JA - JZ
C... Determination of eta of Equation (1)
     MN = JN .AND. 1
```

 C... Commented out by Mark Mattson, April 19, 1996.

```
C
          PN = 1.15
          IF (MZ.EQ.1 .AND. MN.LE.1) PN = 0.9 - 0.1*MN
          IF(MZ.EQ.0 .AND. MN.EQ.0) PN = ET(1)
          IF(MZ.EQ.0 .AND. MN.EQ.1) PN = ET(2)
          IF (MZ.EQ.1 .AND. MN.EQ.0) PN = ET (3)
          IF(MZ.EQ.1 .AND. MN.EQ.1) PN = ET(4)
   C... Determination of cutoff energy, E0 (either 1250 MeV or from Equation (3)
   C... of ref [1].).
         EC = 68.7*AI**0.866
         IF (EC.LT.1250.0) EC = 1250.0
         EI = EJ
         IF (EI.GT.EC) EI = EC
   C... Calculation of correction factor to H(E) in Equation (25) on page 358
   C... of ref [2].
         H3 = 1.0
         IF (EI.LT.80.0) H3 = 1. - EXPF(-(EI/25.)**4)
   C... If the secondary nucleus is higher than helium or if the change in the
   C... number of neutrons is greater than 1, it will be handled later.
         IF(JP.GT.2 .OR. KN.GT.1) GO TO 3
   C... Calculation of correction factor as described on p. 358 of ref [2];
   C... CX .NE. 1 is for light secondary nuclei.
         CX = 1.0
         IF (JZ.EQ.4 .AND. JA.EQ.9) CX = CJ(JA, JZ)/PN
   C... Calculation of correction factor for light nuclei as described on
   C... p. 876 of ref [3].
         IF (AI/ZI.GT.2.0) CT = (ZI - 2.0)/((AI - ZI) - 2.0)
         IF (ZI.GT.5 .OR. CT.GT.1.0 .OR. (AI-ZI).LE.2.0) CT = 1.0
Ŀà
         IF (JP.GT.1) GO TO 2 ! Case of > 1 proton in secondary nucleus.
IF (KN.GT.1) GO TO 3 ! Case of > 1 neutron created in reaction.
Li
C... Cross section for (p,pn) as shown in Equation (24) of ref [2] on p. 357.
         IF (AI.LE.40.0) QN = 24.0*(1.0 + 0.01*AI)
         IF (AI.GT.40.0) QN = 1.02*(AI - 7.0)
         IF (AI.GE.63.0) QN = 57.0
   C... Calculation of H(E) as in Equation (25) on p. 358 of ref [2].
         IF (EI.LT.2500.0)
               FE = (1.0 + 2.1 \times EXP(-(EI/100.0) \times 2) + 0.4 \times EXP(-EI/350.0)) \times H3
   C... Calculation of cross section for (p,pn) reaction.
         QH = QN*CX
         QJ = QH
         IF (EI.LT.EC) QJ = QH*FE
         RETURN
        IF (KN.NE.0) GO TO 3 ! Case for no neutrons created in reaction.
  C... Calculation of cross section for (p,2p) reaction as shown in Equations
  C... (26) and (27) on p. 358 of ref [2].
  C... This is the calculation of H(E) as in Eq. (27) on p. 358 of ref [2].
         IF (EJ.LT.2500.0) FE = (1.0 - EXPF(-(EJ/230.)**2) +
               2.2*EXPF(-EJ/75.0) + 0.33*EXPF(-((EJ-900.0)/500.0)**2))*H3
  C... 21.0 is the value of sigma(E0) as shown in Eq. (26) of ref [2].
```

MZ = JZ .AND. 1

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QH = 21.0*CX*CT

```
QJ = QH*FE
      RETURN
C... If the atomic number is greater than 29, then the cross-section needs
C... to be modified by f(A) and f(E) as outlined in Table 1 on page 340 of
C... ref [2] (FE and FA were previously set equal to 1).
      IF (IZ.LT.29) GO TO 5
C... Calculation of f(E) of Equation (1) of ref [1].
      IF (JZ.NE. 4) FE = (EI/EC)**(0.4*ZJ) ! Helium and Lithium nuclei
      IF (JZ.EQ. 4) FE = (EI/EC)**1.8
                                             ! Beryllium nuclei
C... Calculation of f(A) in Equation (1) or ref [2]; Eq. (10) on p. 351 of
C... ref [2].
      FA = EXPF(0.01*(AI - 56.0)*(AN/ZJ - CN - 0.45))
      IF (FA.LE.1.0) FA = 1.
C...
      EI = EC
      ZI = 29.0
      AI = 63.0
C... Calculation of enhancement factor xi of Equation (1) of ref [1];
C... see Table 2 of ref [3].
      AT = IA
      IF (IA.GE.104) AT = 104
      IF (IA.GE.64) GO TO 7
      IF (IA.GT.34) GO TO 6
C... Situation described in Table 3 of ref [1].
      IF (IA.GE.14 .AND. JA.EQ.6 .AND. JZ.EQ.2) FZ = 1.0 + 0.1*(IA - 14)
      GO TO 8
C... Enhancement factor xi when target nucleus has 34 .LE. IA .LE. 63
      IF (JA.EQ.6 .AND. JZ.EQ.2) FZ = 3.0*(1.0 + 0.02*(IA - 34))
      IF (JA.EQ.6 .AND. JZ.EQ.3) FZ = 1.0 + 0.02*(IA - 34)
      IF (JA.EQ.7 .AND. JZ.LE.4) FZ = 1.0 + 0.01*(IA - 34)
      GO TO 8
C... Enhancement factor xi when target nucleus has 64 .LE. IA .LE. 104
      IF (JA.EQ.6 .AND. JZ.EQ.2) FZ = 4.8 + 0.0450*(AT - 64.0)
      IF (JA.EQ.6 .AND. JZ.EQ.3) FZ = 1.6 + 0.0150*(AT - 64.0)
      IF (JA.EQ.7 .AND. JZ.EQ.3) FZ = 1.3 + 0.0150*(AT - 64.0)
      IF (JA.EQ.7 .AND. JZ.EQ.4) FZ = 1.3 + 0.0105*(AT - 64.0)
      IF (JA.EQ.8 .AND. JZ.EQ.3) FZ = 1.0 + 0.0225*(AT - 64.0)
      IF (JA.EQ.9 .AND. JZ.EQ.3) FZ = 1.0 + 0.0125*(AT - 64.0)
      IF (JA.EQ.9 .AND. JZ.EQ.4) FZ = 1.0 + 0.0100*(AT - 64.0)
      IF (JA.GE.10 .AND. JZ.LE.5) FZ = 1.0 + 0.0050*(AT - 64.0)
      DA = AI - AJ ! Change in atomic weights of target and secondary nuclei.
 8
      AA = DA
C... Calculation of delta A c as in Equation (2) of ref [1].
      AE = 31.5 + 0.052*(AI - 36.0)*(ALOG(EI) - 3.17)
C... Determination of delta A in Equation (1) of ref [1].
      IF (AA.GT.AE) DA = AE
C... Determination of sigma0 in Equation (1) of ref [1].
```

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IF (EI.LT.1250.0) Q1 = QL*EXPF(B*(1.0 - 0.0008*EI))
    C... Determination of P in Equation (1) of ref [1].
          PE = PC
          IF (EI.LT.1250.0) PE = PL*(1. - 0.0008*EI)
   C... Determination of R in Equation (1) of ref [1].
          RE = RC
          IF (EI.LT.1250.0) RE = RL/EI**RU
          IF(IZ.GT.20 .AND. EI.LT.1250.0) RE = 1.8
   C... Correction factor for sigma0 (f3 in Table 1A of ref [1]).
          IF(EI.LT.1250.0 .AND. IZ.GE.21)
                              Q1 = Q1*2.0*EXPF(-((EI - 650.0)/720.0)**2)
        æ
   C... Determination of S in Equation (1) of ref [1].
          IF (AI/ZI.GE.2.0) SS = S - C*(AI/ZI - 2.0)**SE
          IF (AI/ZI.LT.2.0) SS = S + C*(2.0 - AI/ZI)**SE ! See ref [3]
   C... Determination of portion of exponent in Equation (1) of ref [1]
   C... within absolute value symbol.
         ST = (ZJ - (SS - T*AJ)*AJ)
         ZA = (ZJ - (SS - T*AJ)*AJ)**2
   C... Calculation of cross section.
         QR = Q1*EXPF(-PE*DA - RE*ZA)*FA*FZ*CJ(JA, JZ)
         QH = QC*EXPF(-RC*ZA)*FA*FZ*CJ(JA,JZ)
         QE = QH*FE
         IF (IZ.LT.29) QJ = QR
         IF (IZ.GE.29) QJ = QE
C... Enhancement factor for (p,3p) reactions, as shown on p. 876 of ref [3].
         IF (JP.EQ.3 .AND. KN.EQ.0) QJ = 1.5*QR
TH C
         FOLLOWING REVISIONS ARE ADDED 12/27/78
C... From ref [3], the cross section QJ is to be modified by a correction
C... factor.
         IF (IZ.LE.20) RETURN
         QI = QJ
   C... Determination of E' as shown below Equation (3) of ref [3].
         EX = 68.7*56.0**0.866*EJ/EC
         AC = 31.5 + 0.045*(AI - 36.0)*(ALOG(AI) + 1.23)
   C... Equation (3) of ref [3].
         G1 = 1.0 - 0.6*(1.0 - EXPF(-(EJ/1000.0)**2))*EXPF(-(EJ/2000.0)**2)
                + 0.2*(1.0 - EXPF(-(EJ/3000.0)**2))
         GX = 1.0 - 0.6*(1.0 - EXPF(-(EX/1000.0)**2))*EXPF(-(EX/2000.0)**2)
                + 0.2*(1.0 - EXPF(-(EX/3000.0)**2))
   C... Calculation of f2 in Equation (9) of ref [3].
         G2 = 1.0 - 0.4*(1.0 - EXPF(-(EX/2000.0)**2))*
               EXPF(-((EX-1800.0)/1800.0)**2) +
               0.17*(1.0 - EXPF(-(EX/2000.0)**2))
   C... Determination of f1 for the conditions described in Eq. (4) of ref [3].
         IF (EJ.GT.2500.) G1 = GX
         IF (EJ.GT.EC .AND. EJ.LT.2500.0) G1 = SQRT(G1*GX)
```

Q1 = QC

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C... Modification of the cross section.
         IF (IZ.GE.21 .AND. IZ.LE.28) QJ = QR*G1
         IF (IZ.GT.28 .AND. AA.GT.AC) QJ = QE*G2
        RETURN
         END
   C
   С
   С
   C
        SUBROUTINE YIELD2 (IZ, IA, JZ, JA, EJ, QJ)
   C...
   C... This subroutine is for the case where the incoming nucleus is
   C... for elements between boron and sulphur and the secondary nucleus...
   C... is between boron and the incoming nucleus, i.e.,
              5 .LE. IZ .LE. 16 .AND.
                                        5 .LE. JZ .LE IZ
   C...
  C.....
  C... Get parameters is described in Table 1B of ref [1].
        EXTERNAL Y2BLOK
        COMMON /BT/ QM,C5,C6,C7,C8,C9,CD,D1,D2,D3,D4,D5,D6,PC,PL,PU,PG,PH,
S, SE, C, RC, RL, RU, T, ET
        COMMON /FS/ QR,QE,QF,QH,FE,FF,FA,FZ,PJ,PN,GA,ANZJ,AA,AE,AC,EC
  C... Commented out by Mark Mattson, April 19, 1996.
        COMMON /ST/ ST, SS, T
  C... The variable CJ holds the values for the parameter OMEGA of [1].
        DIMENSION CJ(56,26), ET(4), QM(7)
        DIMENSION C5(56), C6(56), C7(56), C8(56), C9(56), CD(56)
<u>į.</u>
        DIMENSION D1 (56), D2 (56), D3 (56), D4 (56), D5 (56), D6 (56)
TU c
        LJ C
        EQUIVALENCE (CJ(1, 9), C9), (CJ(1, 10), CD), (CJ(1, 11), D1), (CJ(1, 12), D2)
₽ c
        EQUIVALENCE (CJ(1,13),D3), (CJ(1,14),D4), (CJ(1,15),D5), (CJ(1,16),D6)
ŭ.
,**[
        DO I=1,56
          CJ(I,5) = C5(I)
          CJ(I,6) = C6(I)
          CJ(I,7) = C7(I)
          CJ(I,8) = C8(I)
          CJ(I,9) = C9(I)
          CJ(I,10) = CD(I)
          CJ(I,11) = D1(I)
          CJ(I,12) = D2(I)
          CJ(I,13) = D3(I)
          CJ(I,14) = D4(I)
          CJ(I,15) = D5(I)
          CJ(I,16) = D6(I)
        ENDDO
        QR = 0.0
        QE = 0.0
        QF = 0.0
        QH = 0.0
        AE = 0.0
```

AC = 0.0

```
FF = 1.0
      FA = 1.0
      FZ = 1.0
      GA = 1.0
      AI = IA
                    ! Real number for the atomic weight of the incoming nucleus
                    į 11
                             11
                                   11
                                       16
      ZI = IZ
                                            11
                                                 number " "
                                                                   11
                  i u
      AJ = JA
                             11
                                   18
                                       11
                                            11
                                                            " secondary
                                                 weight "
                   ! "
                             11
                                   11
      ZJ = JZ
                                                 number "
      AA = AI - AJ ! Difference in atomic weights of nuclei
      AN = AJ - ZJ ! Number of neutrons in the secondary nucleus
      ANZJ = AN/ZJ ! Ratio of neutrons to protons in secondary nucleus
C... Determination of OMEGA in ref [1].
      PJ = CJ(JA, JZ)
      JN = JA - JZ ! Integer for number of neutrons in secondary nucleus
      MN = JN.AND.1 ! Is the # of neutrons even or odd?
      MZ = JZ.AND.1! Is the # of protons even or odd?
      CX = 1.
C... The variable PN is eta as in ref [1].
     PN = 1.15
      IF (MZ.EQ.1 .AND. MN.LE.1) PN = 0.9 - 0.1*MN
      IF(MZ.EQ.0 .AND. MN .EQ.0) PN = ET(1)
      IF (MZ.EQ.0 .AND. MN .EQ.1) PN = ET(2)
      IF(MZ.EQ.1 .AND. MN .EQ.0) PN = ET(3)
      IF(MZ.EQ.1 .AND. MN .EQ.1) PN = ET(4)
C... Determination of E0 ("critical energy") as in ref [1];
C... 1250 MeV is the lower bound.
     EC = 68.7*AI**.866
      IF (EC.LT.1250.0) EC = 1250.0
C...
      EI = EJ
      IF (EI.GT.EC) EI = EC
      H3 = 1.0
      IF (EI.LT.80.0) H3 = 1.0 - EXPF(-(EI/25.)**4)
      KN = (IA - IZ) - (JA - JZ)! Difference in number of n's
                                  ! " " p's (inc. 1st)
      JP = IZ - JZ + 1
      IF((JZ.EQ.7 .AND. JA.EQ.13) .OR. (JZ.EQ.10 .AND. JA.EQ.19))
            CX = CJ(JA, JZ)
C... Correction factor as described on p. 876 of ref [3].
      CT = (ZI - 2.0)/((AI - ZI) - 2.0)
      IF (ZI.GT.5.0 .OR. CT.GT.1.0) CT = 1.0
      IF (JP.GT.1) GO TO 2
      IF (KN.NE.1) GO TO 3
C... Factors for (p, pn) reactions.
C... Calculation of H(E) as in Eq. (25) on p. 358 of ref [2].
      IF (EI.LT.2500.0)
        FE = (1.0 + 2.1 \times EXPF(-(EI/100.0) \times 2) + 0.4 \times EXPF(-EI/350.0)) \times H3
C... Equation (24) on p. 357 of ref [2].
      QH = 24.0*(1.0 + 0.01*AI)*CX
```

FE = 1.0

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OJ = OH
         IF (EI.LT.EC) QJ = QH*FE
        GO TO 10
        IF(JP.GT.2 .OR. KN.NE.0) GO TO 3
  C... Factors for (p, 2p) reactions.
  C... Calculation of H(E) as in Eq. (27) on p. 358 of ref [2].
        IF (EJ.LT.2500.0)
              FE = (1.0 - EXPF(-(EJ/230.0)**2) + 2.2*EXPF(-EJ/75.0)
             + 0.33*EXPF(-((EJ-900.0)/500.0)**2))*H3
  C... Eq. (26) on p. 358 of ref [2] multiplied by correction factors.
         OH = 21.0*CX*CT
        QJ = QH*FE
        GO TO 10
  C... Determination of a portion of sigma0 as in ref [1].
        F1 = QM(4) - QM(5)*ALOG(AI*QM(6))
   C... Determination of the parameter P of [1].
         PE = PC
         IF (EC.GT.1250.0) PE = PG/AI**PH
         IF (EI.LT.EC) PE = PL/EI**PU
C... Determination of the parameter R of [1].
        RE = RC
         IF (EI.LT.1250.0) RE = RL/EI**RU
C... Determination of the parameter S of [1].
         IF (AI/ZI.GE.2.0) SS = S - C*(AI/ZI - 2.0)**SE
         IF (AI/ZI.LT.2.0) SS = S + C*(2.0 - AI/ZI)**SE
  C... Determination of the value in the exponential in Equation (1) in ref [1].
         ST = (ZJ - (SS - T*AJ)*AJ)
         ZA = (ZJ - (SS - T*AJ)*AJ)**2
  C... Determination of sigma0 as in ref [1].
         Q1 = QM(1)*(AI**QM(2) - QM(3))*F1*PE*RE**QM(7)/(1.0-EXPF(-PE*AI))
         QC = QM(1)*(AI**QM(2) - QM(3))*F1*PC*RC**QM(7)/(1.0-EXPF(-PC*AI))
   C... Determination of the cross section.
         QR = Q1*EXPF(-PE*(AI - AJ))*EXPF(-RE*ZA)*CJ(JA,JZ)*PN
         QH = QC*EXPF(-PC*(AI - AJ))*EXPF(-RC*ZA)*CJ(JA,JZ)*PN
         IF(JP.LT.3 .OR. KN.NE.0) GO TO 10
   C...
         QJ = QR*AMIN1(.0022*AJ*AJ,1.)
         IF (JP.NE.4) GO TO 10
         IF (QJ.GT.0.5) QJ = 0.5
   C... Determination of the enhancement factor, xi, as in ref [1].
        IF (IZ.GE.14 .AND. EI.GE.500.0) QJ=QJ*(1.0 + 0.12*(IZ - 13))
         IF (IZ.GE.14 .AND. EI.GE.200.0 .AND. EI.LT.500.0)
             QJ = QJ*(1.0 + 0.12*(IZ-13)*EXP(-((EI-500.0)/350.0)**2))
         IF (JP.EQ.2 .AND. KN.EQ.1 .AND. IZ.GE.12) QJ = QJ*1.7
         IF ((AJ/ZJ).GT.1.8 .OR. IZ.GT.10) RETURN
         IF (JP.NE.1 .OR. KN.NE.2) QJ = QJ*0.3
         IF (JP.EQ.1 .AND. KN.EQ.2) QJ = QJ*0.5
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END
   C
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   C
   C
         SUBROUTINE YIELD3 (IZ, IA, JZ, JA, EJ, QJ)
   C... This subroutine is for the case where the incoming nucleus is
   C... for elements between chlorine and calcium and the secondary
   C... nucleus is between boron and the incoming nucleus, i.e.,
   C...
              17 .LE. IZ .LE. 20
                                 .AND.
                                         5 .LE. JZ .LE. IZ
   C...
   C... Get parameters as in Table 1C of ref [1].
        EXTERNAL Y3BLOK
        COMMON /CT/ BA,QT,P0,P1,P2,P3,P4,P5,R0,R1,R2,S0,S1,T,ET
        COMMON /FS/ QR,QE,QF,QH,FE,FF,FA,FZ,CJ,PN,GA,ANZJ,AA,AE,AC,EC
  C... Commented out by Mark Mattson, April 19, 1996.
        COMMON /ST/ ST,S,T
        DIMENSION
                  BA(92), QT(7), ET(4)
  C... REAL*4 versions of the atomic weights and numbers of the target and
  C... secondary nucleus.
AI = IA
          ZI = IZ
AJ = JA
1
          ZJ = JZ
#
Ž.
          AM = BA(IZ)
IF (AM.EQ.0.0) AM = IA
C... Calculation of E0 as in Equation (3) of ref [1].
          EC = 68.7*AI**0.866
IF (EC.LT.1250.0) EC = 1250.0
EI = EJ
          IF (EI.GT.EC) EI = EC
          QR = 0.0
          QE = 0.0
          QF = 0.0
          QH = 0.0
          AE = 0.0
          AC = 0.0
          DX = 0
          FE = 1.0
          FF = 1.0
          FA = 1.0
          FZ = 1.0
          GA = 1.0
          CJ = 1.0
         AA = AI - AJ
```

RETURN

C... Calculation of last portion of H(E) in Equation (25) of ref [2]. H3 = 1.0 - EXPF(-(AMIN1(EI, 80.0)/25.0)**4)

```
C... Calculation of H4 in Equation (29d) of ref [2].
        H4 = AMIN1 (AMAX1 ((400.0/EI-2.2), 1.0), 2.0)
C... Number of neutrons in secondary nucleus and ratio of neutrons to protons.
        AN = AJ - ZJ
        ANZJ = AN/ZJ
C... Calculation of eta as in Table 1C of ref [1].
        JN = JA - JZ
        MN = JN.AND.1
        MZ = JZ.AND.1
        PN = 1.0
C
С
        IF (MN+MZ.EQ.2) PN = 0.85
C
        IF (MN+MZ.EQ.0) PN = 1.25
C
        IF (MN-MZ.EQ.1) PN = 0.90
        IF(MZ.EQ.0 .AND. MN.EQ.0) PN = ET(1)
        IF(MZ.EQ.0 .AND. MN.EQ.1) PN = ET(2)
        IF(MZ.EQ.1 .AND. MN.EQ.0) PN = ET(3)
        IF(MZ.EQ.1 .AND. MN.EQ.1) PN = ET(4)
C... Total change in number of protons.
        JP = IZ - JZ + 1
C... Total change in number of neutrons.
        KN = IA - IZ - JN
        IF(JP.GT.2 .OR. KN.GT.3 .OR. IA.LT.35) GO TO 14
        IF (JP.EQ.2) GO TO 2
C... Case for where the change in the number of protons is 1, there are no
C... more than two additional neutrons and the target nucleus has an atomic
C... weight greater than 35.
C... Calculation of H(E) as in Equation (25) of ref [2].
        IF (EI.LT.2500.0)
            FE = (1.0 + 2.1/EXPF((EI/100.0)**2) + 0.4*EXPF(-EI/350.0))*H3
C... Calculation of cross section if the weight of the target nucleus is
C... less than or equal to 40, as in Equation (24) of ref [2].
        QH = 24.0*(1.0 + 0.01*AI)
        OJ = OH
C... Determination of variable d as described below Equation (23) in ref [2].
        DX = 3.0
        IF (KN.GT.1) DX = 15.0
        IF (EI.LT.EC) QJ = QH*FE
C... Temporarily finished for (p,pn) reactions.
        IF (KN.EQ.1) GO TO 5
C... (p,pxn) where x .GE. 2. (p. 359 of ref [2]).
        XN = KN
        XA = 1.17
        IF (IZ.LE.30) XA=1.6
C... Calculation of cross section as in Equation (28) of ref [2].
        QH = QH*EXPF(1.0 - XN**(XA - 0.0048*AI))
        QJ = QH
        IF (EI.GE.EC) GO TO 5
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C... Temporarily finished with (p,pxn)
         GO TO 4
 C... (p,2pxn) reactions.
 C... Calculation of f(e) as in Equation (27) of ref [2].
         IF (EJ.LT.2500.0)
             FE = (1.0 - EXPF(-(EJ/230.0)**2) + 2.2/EXPF(EJ/75.0)
           + 0.33/EXPF(((EJ-900.0)/500.0)**2))*H3
C... Calculation of cross section as in Equation (26) of ref [2].
         OH = 21.0
         QJ = QH
         QJ = QH * FE
C... Determination of variable d as described below Equation (23) of ref [2].
         IF (KN.EQ.0) DX = -3.0
         IF (KN.EQ.1) DX = -1.0
C... Temporarily finished with (p,2p) reactions.
         IF (KN.EQ.0) GO TO 5
C... Temporarily finished with (p, 2pxn) where n > 2.
         IF (KN.GT.2) GO TO 14
C... Cross section for (p,2pxn), x .GE. 1 as in Equation (31) of ref [2].
        QH = 17.0
        QJ = QH
        IF (EI.GE.EC) GO TO 5
C... Calculation in change in atomic weights.
        KA = IA - JA
        IF (KA.GE.8) GO TO 5
C... Calculation of (1 + H1)*H3*H4 as almost in Equation (29) of ref [2].
        FE = (1.0 + 1.9/EXP((AA/7.9)**2 + (EI/420.0)**1.4))*H3*H4
        QJ = QH*FE
C... Calculation of Y(IA,IZ) as in Equation (22) in ref [2].
        DD = DX*(AI - AM)/ZI
        YA = EXPF(DD)
        IF (DD.GT.0) YA = 2.0 - 1.0/YA
        IF (IA.LT.35 .OR. IA.GT.209) YA = 1.0
        IF (IA.LT.70 .AND. JP.EQ.3) YA = 1.0
C... Calculation of cross section.
        QH = YA*QH
        QJ = YA*QJ
        RETURN
C... Calculations for when the change in the number of protons is greater
C... than 2, the change in the number of neutrons is greater than 3 or the
C... atomic weight of the target nucleus is less than 35.
C... Calculation of change in atomic weights.
 14
        DA = AI - AJ
C... Determination of OMEGA for what in this region is a special case (see
C... Table 2 of ref [1]).
        IF(JZ.EQ.7 .AND. JA.EQ.13) CJ = 0.39
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F1 = QT(4) - QT(5)*ALOG(AI*QT(6))
    C... Calculation of f2' as in Equation (6) of ref [1].
            IF (EI.GE.600.0) F2 = 1.0
            IF (EI.LT.600.0) F2 = EXPF (0.90 - 0.0015*EI)
            IF (EI.GE.EC) GO TO 15
    C... Determination of P in Table 1C in ref [1] for low energy.
            PE = P0/EI**P1
            IF (EI.LT.EC) GO TO 16
    15
   C... Determination of f2' and P in Table 1C of ref [1] for high energy.
            F2 = 1.00
           PE = P2/AI**P3
   C... Determination of R in Table 1C of ref [1].
           IF (EI.LT.1250.0) RE = R0/EI**R1
           IF (EI.GE.1250.0) RE = R2
   C... Calculation of sigma0 in Table 1C of ref [1].
           Q1 = QT(1)*(AI**QT(2) - QT(3))*F1*F2*PE*RE**QT(7)/
                       (1.0 - EXPF(-PE*AI))
   C... Calculation of S in Table 1C of ref [1].
           S = S0 - S1*ABS(AI/ZI - 2.0)
   C... Determination of value in second exponent of Equation (1) of ref [1].
           ST = (ZJ - (S - 0.0005*AJ)*AJ)
           ZA = ABS(ZJ - (S - 0.0005*AJ)*AJ)**2
   C... Calculation of cross section as in Equation (1) of ref [1].
           QJ = Q1*EXPF (-PE*DA-RE*ZA)
           QR = QJ*PN*CJ
           QJ = QR
C... Consideration of high energy case.
C... Calculation of P in Table 1C of ref [1].
           PH = P4/AI**P5
   C... Determination of R in Table 1C of ref [1].
           RH = R2
   C... Calculation of sigma0 in Table 1C of ref [1].
           QC = QT(1)*(AI**QT(2) - QT(3))*F1*PH*RH**QT(7)/
                      (1.0 - EXPF(-PH*AI))
   C... Calculation of cross section as in Equation (1) of ref [1].
           QH = QC*EXPF(-PH*DA - RH*ZA)*PN*CJ
           IF(JP.NE.3 .OR. KN.GT.0 .OR. IA.LT.35) GO TO 20
   C... Determination of correction factors in (p,3p) reactions.
           FF = 1.0
           FE = 1.0
           QJ = QH
    20
           IF (IZ.LT.14 .OR. IZ.GT.19 .OR. EI.LT.200.0) RETURN
```

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C... Calculation of portion of sigma0 as in Table 1C of ref [1].

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C...
          IF (EI.GT.500.0) QJ = QJ*(1.0 + 0.12*(IZ-13))
          IF (EI.LE.500.0) QJ = QJ*(1.0 + 0.12*(IZ-13)*
              EXP(-((EI-500.0)/350.0)**2))
        RETURN
        END
   C
   C
   C
   С
        SUBROUTINE YIELD4(IZ, IA, JZ, JA, EJ, QJ) ! see notes at end
   C....
   C.....
   C...
   C... This subroutine is for the case where the incoming nucleus is
   C... for elements between scandium and uranium and the secondary
   C... nucleus is between boron and the incoming nucleus, i.e.,
             21 .LE. IZ .LE. 92 .AND. 5 .LE. JZ .LE. IZ
   C...
  C... Get parameters as in Table 1D of ref [1].
        EXTERNAL Y4BLOK
u.
        COMMON /DT/ BA,S0,S1,T3,R0,R1,R2,R3,P0,P1,P2,P3,E0,E1,C1,C2,C3,C4,
                   D1, D2, T2, ET
ř.
        COMMON /FS/ QR,QE,QF,QH,FE,FF,FA,FZ,CJ,PN,GA,ANZJ,AA,AE,AC,EC
        COMMON /Q4/ C0, F1, F2, F3, PE, DA, RJ, ZA, PH, YA, Q1, QM, HM
COMMON /QG/ QI, G1, G2, G3, G4
  C... Commented out by Mark Mattson, April 19, 1996.
#
       COMMON /ST/ ST,S,T2
C... BA is an array representing the average atomic number for all stable
  C... isotopes of a given element.
DIMENSION BA(92), ET(4)
C... REAL*4 versions of atomic weights and numbers of nuclei.
        AI = IA
        ZI = IZ
       AJ = JA
        ZJ = JZ
        AM = BA(IZ) + IA*(300/(INT(BA(IZ)) + 300))
        KM = 0
  C...
        FP = 1.
        QJ = 0.0
        QR = 0.0
        QE = 0.0
        QF = 0.0
        QM = 0.0
       HM = 0.0
       QH = 0.0
       AC = 0.0
       AE = 0.0
       DM = 0.0
```

C... Correction factor xi.

```
FF = 1.0
           FM = 1.0
           FA = 1.0
           FZ = 1.0
           GA = 1.0
           CJ = 1.0
           PN = 1.0
          YA = 1.0
    C... This subroutine doesn't consider reactions where the secondary nucleus
    C... has more mass, protons or neutrons than the target nucleus.
           IF(IA.LE.JA .OR. IZ.LT.JZ .OR. (IA-JA).LT.(IZ-JZ)) RETURN
          AA = AI - AJ
          DA = AA
          AN = AJ - ZJ
          ANZJ = AN/ZJ
          ANZI = AI/ZI - 1.0
          JN = JA - JZ
                                  ! Number of neutrons in product nucleus
                                  ! Change in number of protons
          JP = IZ - JZ + 1
          CN = 0.3*(AI - AM)/ZI
          KN = (IA - IZ) - JN
                                ! Change in number of protons
          XN = KN
    C... Calculation of eta in Table 1D of ref [1].
          MZ = JZ.AND.1
MN = JN.AND.1
    C
          IF ((MN+MZ).EQ.2) PN = 0.85
25.5
   C
          IF ((MN+MZ).EQ.0) PN = 1.25
С
          IF ((MN-MZ).EQ.1) PN = 0.90
譯
          IF (MZ.EQ.0 .AND. MN.EQ.0) PN = ET (1)
IF (MZ.EQ.0 .AND. MN.EQ.1) PN = ET(2)
IF(MZ.EQ.1 .AND. MN.EQ.0) PN = ET(3)
          IF(MZ.EQ.1 .AND. MN.EQ.1) PN = ET(4)
ļ.
    c...
          PN = PN + (1.-PN)*(1.-EXPF(-((IA-100)/35.)**2))
    C... Calculation of DELTA A_c as in Equation (2) of ref [1].
          AC = 31.5 + 0.045*(AI - 36.0)*(ALOG(AI) + 1.23)
          EI = EJ
          EC = E0*AI**E1
          IF (EC.GT.4000.0) EC = 4000.0
          IF (EC.LT.1250.0) EC = 1250.0
          IF (EI.GT.EC) EI = EC
    C... Determination of maximum value for energy dependence of fission cross
    C... section as described on p. 347-8 of ref [2].
          FMAX = 1800.0/EI
          IF (IZ.GE.84) FMAX = (1800.0/EI)**(6.56 - 0.067*ZI)
          IF (FMAX.GT.4.0) FMAX = 4.0
          IF (EI.GT.1800.0) FMAX = 1.0
   C... Determination of OMEGA for certain situations.
          IF (IZ.LE.28 .AND. JZ.EQ.20 .AND. JA.EQ.19) CJ = 0.6
          IF (JZ.EQ.7 .AND. JA.EQ.13) CJ = 0.39
```

DX = 0.0 KFLAG = 0 FE = 1.0

```
C... Determination of parameter P of Table 1D of ref [1] for different
    C... energies.
          P500 = P2/(500.0**P3)
          P1000 = P2/(1000.0**P3)
          P3000 = P2/(3000.0**P3)
          IF (JP.GE.6 .OR. IA.LT.35) GO TO 100
    C...
          IF (JP.GE.4.AND.JP.LE.5.AND.KN.GE.KM.AND.EJ.LT.EC) GO TO 100
          PERIPHERAL
    C... Calculation of H3 as in Equation (29c) of ref [2].
          H3 = 1.0 - EXPF(-(EI/(15.0 + 10.0*AA))**4)
   C... Determination of H4 as in Equation (29d) of ref [2].
         H4 = 400.0/EI - 2.2
          IF (H4.LT.1.0) H4 = 1.0
          IF (H4.GT.2.0) H4 = 2.0
   C... Calculation of H(E) as in Equation (29) of ref [2].
         HE = (1.0 + 1.9 \times EXPF(-(DA/7.9) \times 2 - (EI/420.0) \times 1.4)
                 - (1.0 - EXPF(-(DA/12.0)**8))*EXPF(-(EI/420.0)**3))*H3*H4
         EC = 2500.0
         IF (JP.GE.3 .AND. JP.LE.5) GO TO 3
         IF (JP.EQ.2) GO TO 2
C... For (p,pn) reactions.
\square C... Calculation of H(E) as in Equation (25) of ref [2].
         IF (EI.LT.2500.0)
             FE = (1.0 + 2.1*EXPF(-(EI/100.0)**2) + 0.4*EXPF(-EI/350.0))*H3
         IF (KN.GT.1) FE = HE
C... Calculation of x_max as in Equation (30) of ref [2].
         KM = AI/20.0 + 1.5*(ABS(238.0 - AI)/167.0)**2.5 + 0.8
         IF (IA.LE.70) KM = 3.0 + AI/66.0
         IF (KN.GT.KM) GO TO 100
   C... Calculation of cross section for (p,pn) as in Equation (24) of ref [2].
         IF (IA.LE.40) QA = 24.0*(1.0 + 0.01*AI)
         QH = QA
         IF (IA.GT.40) QA = 1.02*(AI - 7.0)
         QH = QA
         IF (IA.GE.63) QA = 57.0
         QH = QA
  C... Determination of exponent of x in Equation (28) of ref [2] (see also
  C... p. 876 of ref [3]).
        XA = 1.17
        IF (IZ.LE.30) XA = 1.5
  C... Calculation of cross section for (p,pxn) as in Equation (28) of ref [2].
        IF (KN.NE.1) QH = QA*EXPF(1.0 - XN**(XA - 0.0048*AI))
  C... Determination of d as described below Equation (23) of ref [2].
        IF (KN.EQ.1) DX = 3.0
        IF (KN.GT.1) DX = 15.0
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QH = QH*(1. + 0.15*(IZ/80.)**2)
         GO TO 6
   C... Calculation of H(E) as in Equation (27) of ref [2].
         FE = (1.0 - EXPF(-(EJ/230.0)**2) + 2.2*EXPF(-EJ/75.0)
             + 0.33*EXPF(-((EJ-900.0)/500.0)**2))*H3
         IF (KN.GT.0) FE = HE
   C... Determination of x max as in Equation (32) of ref [2].
         KM = 10.0*(1.0 - EXPF(-((AI-39.0)/54.5)**2))
         IF (KM.LT.2) KM = 2
         IF (KN.GT.KM) GO TO 100
   C... Calculation of cross section as in Equation (31) of ref [2].
         OH = 17.0
         IF (KN.GT.0 .AND. IZ.LE.50) QH = 27
         IF (KN.GT.0 .AND. IZ.GT.50) QH = 21
         IF (KN.EQ.0 .AND. IZ.LE.28) QH = 33
         IF (KN.EQ.0 .AND. IZ.LE.23) QH = 28
         IF (KN.EQ.0 .AND. IZ.LE.5) QH = 21
         QJ = QH
C... Determination of d as described below Equation (23) of ref [2].
         IF (KN.EQ.0) DX = -3.0
         IF (KN.EQ.1) DX = -1.0
         GO TO 6
   C... Calculation of H*(E) as below Equation (34a) in ref [2].
   C...
lest
         FN = AMIN1((EI/4200.)**(0.72 - 0.18*XN)*H3,1.0)
IF (IA.LE.70) FE = 1.0 - EXPF(-(EI/35.0)**4)
         IF (IA.GT.70 .AND. KN.LE.4) FE = FN
į.
         IF (IA.GT.70 .AND. KN.GT.4) FE = H3
   C... Calculation of H(E) as in Equation (34a) of ref [2].
         IF (EI.LT.200.0) FE = FE*EI/200.0
   C... Calculation of x max as in Equation (35) of ref [2].
         KM = AI/25.0 + 0.5
         XM = KM
         IF (KN.GT.KM) KFLAG = 1
         IF (IA.LE.70.0) KFLAG = 0
   C... Calculation of H(E) as in Equation (34a) of ref [2].
         FM = FE
         IF (KN.LE.4) FM = (EI/420.0)**(0.72-0.18*XM)*H3
         IF (FM.GT.1.0) FM = 1.0
         IF (EI.LT.200.0) FM = FM*EI/200.0
   C... Calculation of cross section as in Equation (33b) of ref [2].
         QH = 0.2 + 60.0*(EXPF(-((AI - 89.0)/25.0)**2 - ((XN-4.6)/2.0)**2)
               + (1.0 - EXPF(-(AI/135.0)**3))*
               EXPF(-((XN - AI**0.46)/AI**0.27)**2))
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HM = 0.2 + 60.0*(EXPF(-((AI-89.0)/25.0)**2-((XM-4.6)/2.0)**2)

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+ (1.0 - EXPF(-(AI/135.0)**3))*
             EXPF(-((XM - AI**0.46)/AI**0.27)**2))
C... Determination of d as described below Equation (23) of ref [2].
       IF (KN.EQ.0) DX = -10.0
       IF (KN.EQ.1) DX = -3.0
       IF (IA.LE.70) GO TO 100
       IF (KN.LE.2) QH = (1.2E5/AJ**2.2)*0.12*EXPF(XN/0.85)
       IF (KN.GE.3 .AND. IZ.LE.26) QH = (1.2E5/AJ**2.2)*EXPF((XN-2)/2.0)
      IF ((JZ.LT.82 .AND. KN.GT.KM) .OR. (IZ.GE.88 .AND. JP.GT.3))
                                   KFLAG = 1
       QH = 0.1**(JP - 3)*QH
       HM = 0.1**(JP - 3)*HM
      IF (EI.GE.EC) FE = 1.0
C... Calculation of Y(A, Z) as in Equation (22) of ref [2] and Equation (3)
C... of ref [3].
       IF ((DX*(AI-AM)).LE.O) YA = EXPF(DX*(AI - AM)/ZI)
       IF ((DX*(AI-AM)).GT.0) YA = 2.0 - EXPF(-DX*(AI-AM)/ZI)
       IF (JP.GE.3 .AND. IA.LE.70) YA = 1.0
C... Y(A,Z) includes correction factor phi(A,E) as in Equation (38)
C... of ref [2].
     IF (IA.GT.157 .AND. EI.GT.500 .AND. (JP*KN).NE.1)
             YA = YA*(1.0 - 0.012*(AI-157.0)*(1.0 -
     &
             EXPF(-((EI-500.0)/290.0)**2)))
             *(1.0 - EXPF(-0.6*(1.0*JP)**0.8-((92.0-ZI)/2.7)**2))
C... A product with a nucleus at least as large as carbon's or if it's at
C... least as large as beryllium's with an incoming nucleus greater than 88,
C... we'll handle it later.
       IF (JP.GT.5 .OR. (JP.GT.3 .AND. IZ.GE.88)) GO TO 100
      QH = QH*YA
C... Calculation of cross section as in Equation (37) of ref [2].
       IF (IZ.EQ.92 .AND. JZ.EQ.88) QH = 1.2*EXPF(-0.70*ABS(JA - 224.0))
       IF (IZ.EQ.92 .AND. JZ.EQ.89) QH = 1.6*EXPF(-0.15*ABS(JA - 224.0))
       IF (IZ.EQ.92 .AND. JZ.EQ.90) QH = 8.0*EXPF(-0.25*ABS(JA - 233.0))
       IF (IZ.EQ.91 .AND. JZ.EQ.91) QH = 18.5*EXPF(-0.55*ABS(JA - 234.0))
       IF (IZ.EQ.92 .AND. JZ.EQ.9) QH = 55.0*EXPF(-0.80*ABS(JA - 237.0))
       QJ = QH*FE
      HM = HM*YA
      QM = HM*FM
       IF (KFLAG.EQ.1) GO TO 100
      GO TO 200
C... Calculation of E0 as in Equation (3) of ref [1].
 100 EC = E0*AI**E1
       IF (EC.GT.4000.0) EC = 4000.0
C... (see end of subroutine)
       IF (IA.GT.100) FP = 1. - 0.20*(IA-100)/100.
       IF (IA.GT.180) FP = 0.84
      ANZX = AMIN1(ANZJ,ANZI)
      ANZC = ANZJ - CN
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AZ = ANZJ
          EE = 450.0/EI
   C... Calculation of f_FU(E) as in Equation (20a) of ref [2].
          IF (JA.GT.36 .AND. JZ.LT.88 .AND. IZ.EQ.92)
                FF = EXPF(33.0*((ANZJ - 1.36) + 0.00006*EI-1200.0/
                     (EI+80.0) **2))
         IF (JA.GT.36 .AND. JZ.LT.88 .AND. IZ.EQ.90)
                FF = EXPF(16.0*EE**0.6*((ANZJ - CN) + 0.00006*EI -
                1200.0/(EI + 80)**2))
         IF (JA.GE.36 .AND. JZ.LT.IZ .AND. IZ.GE.84 .AND. IZ.LT.90)
                FF = EXPF(19.0*EE**0.6*((ANZX - CN) - ALOG(5.5/EI**0.07)))
   C... Determination of g as on p. 349 of ref [2].
         GE = EE
         IF (EI.LE.200.0) GE = 2.25
   C... Determination of h as on p. 349 of ref [2].
         H1 = AJ - 105.0 + 0.68*(207.0 - AI)
         IF (EI.GT.450.0 .AND. EI.LT.600.0) H1 = 0.0006*H1*(600.0 - EI)
         IF (EI.LE.450.0) H1 = 0.0900*H1
         IF (EI.GE.600.0 .OR. AI.LT.180.0 .OR. H1.LT.0) H1 = 0.0
   C... Determination of portion of f_F(E) as shown in Equation (13) of ref [3].
         AE = 0.0065*(207 - IA)*EXPF(-ABS(EI - 700.0)/700.0)
   C... Calculation of f_F(E) in Equation (5) of ref [2].
         IF (IA.GE.110 .AND. IZ.LE.83 .AND. JZ.LT.IZ)
        &
             FF = EXPF(19.0*EE**0.6*((ANZX - CN) - ALOG(5.5/EI**0.07) - AE)
               -(((AJ/AI - 0.46)/0.15)**2*GE+H1))
        IF (IA.GE.110 .AND. IZ.LE.83 .AND. JZ.LT.IZ .AND.
            (ANZX - CN).LT.1.3) FF =
              FF*EXPF((1.3-ANZX + CN)*(350.0/EI)**4/(1.0 + (130.0/EI)**4))
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         IF (FF.GT.FMAX) FF = FMAX
\mathbb{L} C... Determination of f_B(E) as in Equation (14) of ref [2].
Lâ
         IF (JZ.GE.88) GO TO 110
         EE = EI/EC
         A4 = (AI/63.0)**4
٠. أيب<sup>يد</sup>
         IF (EE.GE.0.3) FE = EE**(2.2 - 0.01*A4)
         IF (EE.LT.0.3) FE = EE**(2.2 + 0.01*A4)*0.3**(-0.02*A4)
         IF (JZ.GE.5 .AND. JZ.LE.8) FE = EE**1.8
         IF (JZ.GE.9 .AND. JZ.LE.10) FE = EE**2.0
         GF = 1.0
         EX = EI
         IF (EX .LT. 500.0) EX = 500.0
  C... Determination of G as in Equation (15) of ref [2].
         IF(JA.GE.18 .AND. JA.LE.30)
               GF = EXPF(2600.0*((ANZX - CN) + 0.01*AI - 3.1)/EX)
        IF(JA.GE.31 .AND. JA.LE.56)
               GF = EXPF(5600.0*((ANZX - CN) + 0.0035*AI - 1.72)/EX)
        IF(GF.LT.1.0) GF = 1.0
  C... f_B(E) as in Equation (14) of ref [2].
        FE = FE*GF
        IF(JZ.LT.30 .OR. IZ.LE.83) GO TO 140
  C
  C
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IF (ANZC.GT.1.5) ANZC = 1.5

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C... As the various references suggest that data for very large nuclei is
C... sparse at best, I have no idea where these next few lines came from,
C... save that they are used to modify the parameter eta (called CJ).
 110 	ext{ FU} = 1.0
      IF (IZ.GE.88 .AND. JZ.GE.82 .AND. (KN.GT.KM .OR. JP.GT.3))
             FU = (1.0 - EXPF(-0.6*(1.0*JP)**0.8 - ((92.0 - ZI)/2.7)**2))
             *(1.0 - 0.8*(EXPF(-((ZI - ZJ)/5.4)**8)))
      IF (IZ.GE.88 .AND. JZ.GE.57 .AND. JZ.LT.82) FU =
            1.0 - 0.36*EXPF(-((82.0 - ZJ)/27.0)**2 - ((92.0-ZI)/2.7)**2)
      CJ = CJ*FU
C... Calculation of cross section as per Equation (18) of ref [2].
 120 IF(JZ.GE.30 .AND. JZ.LE.35)
             QH = 6.5 \times EXPF(-80.0 \times (AN/ZJ - 1.26) \times 2) \times PN
      IF(JZ.GE.36 .AND. JZ.LE.43)
             QH = 10.0 \times EXPF(-55.0 \times (AN/ZJ - 1.33) \times 2) \times PN
      IF(JZ.GE.44 .AND. JZ.LE.50)
             QH = 10.0 \times EXPF(-70.0 \times (AN/ZJ - 1.36) \times 2) \times PN
      IF (JZ.LE.50) GO TO 130
C... This is mostly described in Equation (18d) of ref. [2] including the
C... discussion following the equation about extrapolation into regions of
C... nuclei larger than Z > 55.
      EE = EI - 100
      IF (EE.LT.150.0) EE = 150.0
      HE = 0.03*(ALOG(EE))**2
                                                 ! Equation (20b) of ref [2].
      IF (HE.GT.1.0) HE = 1.0
      ZZ = JZ
      IF (JZ.LT.55) ZZ = 55.0
      HS = (1.5 \times EXPF(-1.0/ZJ \times (ZZ - 55.0) \times 2) + 3.0) \times FU \times HE
      HF = 5.0 \times EXPF(-10.0/ZJ \times (ZZ - 55.0) \times 1.5)
      CON = 80.0 + 220.0*(1.0 - EXPF(-(ZJ - 55.0)**2/4.0))*(JZ/55)
      IF(JZ.GE.51 .AND. JA.LE.155)
            QH = HF*EXPF(-CON*(AN/ZJ - 1.48)**2)*PN
     &
             + HS*EXPF(-260.0*(AN/ZJ - 1.29)**2)*PN
      IF(JZ.GE.66 .AND. JZ.LE.87) GO TO 140
      IF (IZ.LT.88) GO TO 140
      QJ = QH
C... Correction for Z = 89 and Z = 90; see p. 353 of ref [2].
      IF (IZ.LE.90) QH = 0.6*QH
 130 QF = QH\starFF
      QJ = QF
      GO TO 200
C... Calculation of f(A) as in Equations (10) and (13) of ref. [2] (see also
C... discussion in section (f) of ref [3]).
 140 IF (IZ.GE.29 .AND. JZ.EQ.5) FA = EXPF(0.02*(AI - 56)*(ANZC - 0.6))
      IF (IZ.GE.29 .AND. JZ.GE.6 .AND. JZ.LE.8)
             FA = EXPF (0.020*(AI - 56.0)*(ANZC-0.7))
      IF (IZ.GE.29 .AND. JZ.GE.9)
            FA = EXPF(0.020*(AI - 56.0)*(ANZC-0.9))
      IF (JZ.LE.11) FA = FA*EXPF(-2.5*(ANZC - 1.0))
      IF (FA.LT.1.0) FA = 1.0
C... See Table 1 of ref [2], which determines regions of applicability for
C... the various ways of calculating the cross section.
      IF (IA.GE.110) FM = FF
```

```
FM = AMAX1(FE, FF)
          IF(IA.GE.110 .AND. JA.LT.57 .AND. AJ.LE.(0.23*AI)) FM = FE
    C... Determination of Delta A c as in Equation (2) of ref [1].
          DMAX = 31.5 + 0.045*(AI - 36.0)*(ALOG(AI) + 1.23)
          AC = DMAX
          AE = AC
          AH = DA
          IF (AH.GT.DMAX) AH = DMAX
          IF (EI.LT.EC) DMAX = 31.5 + D1*(AI - 36)*(ALOG(EI) - D2)
          AE = DMAX
          IF (DA.GT.DMAX) DA = DMAX
   C... Calculation of fl as in Equation (4) of ref [1].
    150 F1 = EXPF(-0.25 + 0.0074*AI)
          IF (EI.GT.750.0) F1 = 1.0 + (F1 - 1.0)*((EC - EI)/(EC - 750.0))**2
          IF (EI.GE.EC) F1 = 1.0
   C... Calculation of f2 as in Equation (5) of ref [1].
          F2 = EXPF(1.73 - 0.0071*EI)
          IF (F2.LT.1.0) F2 = 1.0
   C... Calculation of parameter P in Table 1D of ref [1].
          PE = P2/EI**P3
          PH = PO/AI**P1
   C... Calculation of Cp as below Table 1D of ref [1].
          IF (IZ.GE.20 .AND. IZ.LE.30)
                PE = PE*(1.0 - 0.32*EXPF(-((EI - 100.0)/100.0)**2))
         &
Li
         IF (IA.GT.100)
Li
           PE = PE*(1.0 - 0.000015*(AI - 100.0)*(EC + 150.0)/(EI + 150.0))
in all
         IF (IA.LE.71 .AND. EJ.GE.EC) PE = PH
IF (IA.GT.72 .AND. EJ.GT.3000.0) PE = PH
L
         IF (EJ.GE.EC.AND.EJ.GE.3000.0) PE = PH
.
Essêt
         PX = 0.0980*(1000.0/EJ)**(0.819*ALOG(AI) - 2.732)
1
         IF (EJ.GT.1000.0 .AND. EJ.LT.3000.0) PE = PX
         PA = PE * AI
         HA = PH * AI
   C... Determination of parameter R of Table 1D of ref [1].
         RJ = R0*AJ**(-R1)*(1.0 - 0.4*(IZ/88))
         IF (AJ.LT.40.0) RJ = R2*AJ**R3*(1.0 - 0.4*(IZ/88))
   C... Calculation of A' as discussed below Equation (2a) of ref [2].
          IF (AA.LT.AC .AND. JZ.EQ.40) DM = -1.1
          IF (AA.LT.AC.AND.JZ.EQ.42) DM = 0.8
   C...
         IF (AA.LT.AC .AND. JZ.EQ.53) DM =-0.5
         IF (AA.LT.AC .AND. JZ.EQ.55) DM = 1.7
         IF (AA.LT.AC.AND.JZ.EQ.60) DM = -1.1
         IF (AA.LT.AC .AND. JZ.EQ.61) DM = -1.3
         IF (AA.LT.AC.AND.JZ.EQ.63) DM = 0.9
         IF (AA.LT.AC.AND.JZ.EQ.64) DM = 0.7
```

IF (IZ.GT.76 .AND. JZ.GT.62) DM = DM + 1.0

AJ = AJ + DM

IF(IA.GE.110 .AND. JA.LT.57 .AND. AJ.GT.(0.23*AI))

IF (IA.LT.110) FM = FE

```
C... Determination of parameter S in Table 1D of ref [1].
         S = S0 - S1*(AI - AM)/ZI
   C... This modified version of the atomic number of the secondary nucleus
   C... doesn't seem to have a counterpart description in any of the papers.
         ST = ZJ - (S - T2*AJ - T3*AJ*AJ)*AJ
         IF (ST.LT.-1.0) ZA = ABS (ST)**1.30
         IF (ST.GE.-1.0) ZA = ABS (ST)**1.50
         IF (ST.GT. 1.0) ZA = ABS(ST)**1.75
   C... Calculation of cross section as in Equation (1) of ref [1].
         Q0 = C1*F1*F2*PE*AI**C2/(1.0 - C3/PA - (C4 - C3/PA)*EXPF(-PA))
         HO = C1*1.0*PH*AI**C2/(1.0 - C3/HA - (C4 - C3/HA)*EXPF(-HA))
         QR = CJ*PN*EXPF(-PE*DA - RJ*ZA)*Q0
         QH = CJ*PN*EXPF(-PH*AH - RJ*ZA)*H0
   C... Evaluation of A0 as discussed in Equation (16) of ref [3].
         AX = 0.5*(S - SQRT(S*S - 4.0*T2*ZJ))/T2 - DM
   C... Evaluation of x0 somewhat as in Equation (15) of ref [3].
         KX = (AI - AX) - (JP - 1)
         XP = KX
         IF (IA.GT.70 .AND. KN.GT.KM .AND. KN.LT.KX .AND. KFLAG.EQ.1)
                             GO TO 20
         GO TO 195
   C... Calculation of f1 as in Equation (4) of ref [1].
        F1000 = 1.0 + (EXPF(0.0074*AI - 0.25) - 1.0)*
               ((EC-1000.0)/(EC - 750.0))**2
C... Calculation of cross section as in Table 1D and Equation (1) of ref [2].
        Q1000 = CJ*PN*C1*F1000*AI**C2*P1000*EXPF(-P1000*DA-RJ*ZA)
               /(1.0 - C3/P1000/AI - (C4 - C3/P1000/AI)*EXPF(-P1000*AI))
C... Cross section calculated with different DELTA A.
        X1000 = Q1000*EXPF(-P1000*(AI - AX))/EXPF(-P1000*DA - RJ*ZA)
C... Yet another correction to the cross section.
        HR = QH*EXPF(-PH*(AI - AX))/EXPF(-PH*AH - RJ*ZA)
        IF (X1000.GT.HR) X1000 = HR
  C... Repeat above procedure to calculate different cross section, but for
  C... a different energy.
        F500 = EXPF(0.0074*AI - 0.25)
        Q500 = CJ*PN*C1*F500*AI**C2*P500*EXPF(-P500*DA - RJ*ZA)
              /(1.0 - C3/P500/AI - (C4 - C3/P500/AI) *EXPF(-P500*AI))
        X500 = Q500*EXPF(-P500*(AI - AX))/EXPF(-P500*DA - RJ*ZA)
        QR = QR*EXPF(-PE*(AI - AX))/EXPF(-PE*DA - RJ*ZA)
        IF (EI.GT.EC) QR = HR
        IF (EI.GE.1000.0 .AND. QR.GT.HR) QR = HR
        IF (EI.LT.1000.0 .AND. EI.GT.500.0)
              QR = X500 + (X1000 - X500)*(EI - 500.0)/500.0
        QH = HM + (HR - HM) * (XN - XM) / (XP - XM)
        QJ = QM + (QR - QM) * (XN - XM) / (XP - XM)
        KFLAG = 0
        GO TO 200
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KFLAG = 0

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IF (EJ.GT.EC) QR = QH

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IF (EI.GE.3000.0 .AND. QR.GT.QH) QR = QH
   C... Repeat procedure described above, but for different circumstances.
          F500 = EXPF(0.0074*AI - 0.25)
         Q500 = CJ*PN*C1*F500*AI**C2*P500*EXPF(-P500*DA - RJ*ZA)
               /(1.0 - C3/P500/AI - (C4 - C3/P500/AI)*EXPF(-P500*AI))
         F1000 = 1.0 + (EXPF(0.0074*AI - 0.25) - 1.0)*((EC-1.0E3)
               /(EC-750.0))**2
         01000 = CJ*PN*C1*F1000*AI**C2*P1000*EXPF(-P1000*DA - RJ*ZA)
               /(1.0 - C3/P1000/AI - (C4 - C3/P1000/AI)*EXPF(-P1000*AI))
         F3000 = 1.0 + (EXPF(0.0074*AI - 0.25) - 1.0)*
               ((EC - 3.0E3)/(EC -750.0))**2
         Q3000 = CJ*PN*C1*F3000*AI**C2*P3000*EXPF(-P3000*DA - RJ*ZA)
               /(1.0 - C3/P3000/AI - (C4 - C3/P3000/AI)*EXPF(-P3000*AI))
         FDAE = AMIN1((AI - AJ)/(0.14*AI)*(EI/1000.0)**(-2.0/3.0),2.0)
         IF (IZ.GE.29 .AND. IZ.LE.83 .AND. EI.LE.1.0E3) QR = QR*FDAE
         IF (IZ.GE.29 .AND. IZ.LE.83 .AND. EI.GT.1.E3 .AND. EI.LE.3.0E3)
               QR = 0.5*(3.0*Q1000 - Q3000) + (Q3000 - Q1000)/2000.0*EI
         QJ = QR
         QI = QR
         IF (IZ.LT.88 .OR. JP.GT.3) GO TO 160
         QJ = QR*FU
         GO TO 200
    160 DD = AC - AA
   C... Calculation of (N/Z)c as in Equation (3) of ref [2].
         XZ = 1.29 + 0.005*DD + CN
         IF (OH.GT.0.0) RH = OR/OH
         IF (GF.GT.0.0) FR = FE/GF
C... Calculations for different regions as discussed in Table 1 of ref [2].
         IF (IA.GE.69 .AND. RH.LT.FR .AND. EI.LT.300.0) QR = QH*FR
         IF (IZ.GE.76 .AND. IZ.LE.80) FF = FF*AMIN1(1050.0/EI + EI/EC.6.0)
         IF (IA.GE.110 .AND. IA.LE.238) QF = QH*FF*FA
         IF (IA.GE.69 .AND. JA.LE.57) QE = QH*FE*FA
         IF (IA.LT.110. OR. IA.GT.209 .OR. JA.LE.56) GO TO 190
         IF (DD.GE.20) GO TO 170
         IF (IA.LE.125) GO TO 180
         IF (AZ.LE.XZ .OR. JZ.GT.57) GO TO 170
         KZ = 51 + IZ/76 + IZ/80
         IF (AZ.GT.XZ .AND. JZ.LT.KZ) GO TO 180
   C... Calculation of exponent gamma as described below Equation (7) of ref [2]
   C... and in Equation (19) of ref [3].
         ER = EI**(2.0/3.0)
         IF (ANZC.LT.1.56) GA = (0.03 + (ZJ - KZ)*0.007)*(1.56 - ANZC)*ER
         IF (ANZC.GE.1.56) GA = 0
         IF (GA.GT.1.0) GA = 1.0
   C... Calculation of cross section as in Equation (7) of ref [2].
         IF (JZ.GE.KZ .AND. JZ.LE.57 .AND. DD.LT.20.0 .AND. AZ.GT.XZ)
               QJ = QR**GA*QF**(1.0 - GA)
         GO TO 200
   C... Cross sections for different regions as discussed in Table 1 of ref [2].
    170 	 QJ = QR
```

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GO TO 200
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180 QJ = AMAX1(QR, QF)
          GO TO 200
   C... More regions discussed in Table 1 of ref [2].
    190 IF(IA.GE.210 .AND. JA.GE.57) QJ = QF
         IF(IA.GE.110 .AND. JA.LE.56 .AND. AJ.GT.(0.23*AI)) QJ = QH*FA*FM
         IF(IA.GE.110 .AND. JA.LE.56 .AND. AJ.LE.(0.23*AI)) QJ = QE
         IF(IA.LT.110 .AND. IA.GE.69 .AND. DD.GE.0) QJ = QR
         IF(IA.LT.110 .AND. IA.GE.69 .AND. DD.LT.0) QJ = QE
         IF(IA.LT.69) QJ = QR
   C... Situation described in Equation (18d) and below in ref [2].
         IF (QJ.EQ.QR .AND. QI.GT.0.0) QH = QH*QR/QI + 0.000001
         IF (IZ.GE.90 .AND. JZ.GE.66)
              QJ = HF*EXPF(-CON*(AN/JZ - 1.48)**2)*PN*(0.6 + 0.2*(IZ - 90))
               + HS*EXPF(-RJ*ZA)*PN*(0.6 + 0.2*(IZ - 90))
   C... Determination of exponent gamma.
         IF (QJ.EQ.QE .OR. QJ.EQ.QF) GA = 0.0
         IF (QJ.EQ.QR) GA = 1.0
         IF (JP.LT.3 .OR. JP.GT.5) GO TO 200
\square C... Determination of H*(E) as in Equations (34a) and (34b) of ref [2].
         FE = 1.0 - EXPF(-(EI/35.0)**4)
         IF (EI .LT. 200.0) FE = FE*EI/200.0
         IF (JP.GE.3 .AND. KN.GE.1 .AND. IA.LE.70) GO TO 200
  C...
         IF (IZ.GT.30.AND.JP.GE.3.AND.KN.GE.KM) GO TO 200
        QJ = QH*FE*YA
  C... Calculation of E' as used in ref [3];
  C... justification of equation not found, however.
   200 EX = E0*56.0**E1*EJ/EC
  C... Determination of correction factor fl as in Equation (3) of ref [3].
        G1 = 1.0 - 0.6*(1.0 - EXPF(-(EX/1000.)**2))*EXPF(-(EX/2000.0)**2)
              + 0.2*(1.0 - EXPF(-(EX/3000.0)**2))
  C... Determination of correction factor f2 as in Equation (9) of ref [3].
        G2 = 1.0 - 0.4*(1.0 - EXPF(-(EX/2000.0)**2))*
              EXPF(-((EX - 1800.0)/1800.0)**2)
              + 0.17*(1.0 - EXPF(-(EX/2000.0)**2))
  C... Determination of correction factor f3 as in Equation (10) of ref [3].
        G3 = 1.0 + 0.25*(1.0 - EXPF(-(EX/1500.0)**2))*
              EXPF(-((EX - 1500.0)/1800.0)**2)
              - 0.05*(1.0 - EXPF(-(EX/2000.0)**2))
  C... Determination of correction factor f4 as in Equation (11) of ref [3].
        G4 = 1.0 - 0.1*(1.0 - EXPF(-(EX/4000.0)**2))
  C... Application of correction factors as described in part (b) of ref [3].
        IF (IZ.LE.28 .AND. JZ.LE.4 .AND. JA.LE.12) QJ = QJ*G1
        IF (IZ.LE.28 .AND. AA.GT.AC) QJ = QJ*G2
```

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IF (IZ.GE.90 .AND. JA.GT.56 .AND. AA.GE.7.0) QJ = QJ*G4
   C... Correction for when the change in the number of neutrons is much larger
   C... than the change in the number of protons.
        MX = 10
        IF (IA.LT.150 .AND. JP.LT.3) MX = 9
        IF (IA.GE.150 .AND. JP.EQ.3) MX = 11
        IF (JP.EQ.1 .AND. KN.GT.MX) QJ =
                                   QJ*(0.1 +0.9*EXPF(-(XN - MX)**2/4.0))
        IF (JP.EQ.2 .AND. KN.GE.MX) QJ =
                                   QJ*(0.1 +0.9*EXPF(-(XN - MX)**2/4.0))
        IF (JP.EQ.3 .AND. KN.GE.MX) QJ =
                                   QJ*(0.5 + 0.5*EXPF(-(XN - MX)**2/4.0))
  C...
        QJ = QJ*FP
        RETURN
        END
  C
  C
  C
  С
        SUBROUTINE PXN(Z, A, X, E, QJ)
  C.....
  C.....
  C... This subroutine is for the case where the incoming nucleus is
  C... for elements between scandium and uranium and the secondary
  C... nucleus is between boron and the incoming nucleus, i.e.,
  C...
             21 .LE. IZ .LE. 92 .AND. 5 .LE. JZ .LE. IZ
=
  C...
        COMMON /A/ Q1, Q2, QI, F1, F2, F3
EI = 500.0 + 300.0*X
Ü
        Q1 = 3.3 \times EXPF(-ABS(6.9 - X) \times 2.8/67.45 -
                 90.0*X**2.35*ABS(2.5 - A/Z)**5)
        QA = 3.3*EXPF(-ABS(6.9 - X)**2.8/67.45 - 0.04638*X**2.35)
        QB = 3.3*EXPF(-ABS(6.9 - X)**2.8/67.45)
        IF (A/Z.GE.2.28 .AND. A/Z.LE.2.50)
              Q1 = QA**((2.5 - A/Z)/0.22) * QB**((A/Z - 2.28)/0.22)
        IF (A/Z.GT.2.5) Q1 = QB
        QI = Q1
        XZ = ((A - Z) - X + 1)/Z
        Q2 = 0.5 \times EXPF(-90.0 \times ABS(1.5 - XZ) \times 5)
        IF (XZ.LE.1.2) QI = AMIN1(Q1, Q2)
        IF (Z.GE.81.0) FI = 1.5*EXPF(-X*((Z - 80.)/12.0)**5)
        IF (FI.GT.1.0) FI = 1.0
        IF (Z.GE.81.0) QI = QI*FI
        IZ = Z + 0.1
        NX = X + 0.1
        IA = A + 0.1
        NT = IA - IZ
        N = NT - NX
        ED = E
```

IF (NX.GE.3 .AND. IZ.GE.39 .AND. NT.GE.28 .AND. N.LT.50)

ED = E - 10.0

IF (IZ.GT.28 .AND. JA.LE.56 .AND. AA.GT.AC) QJ = QJ*G2

IF (AA.GE.7.0 .AND. AA.LE.(AC-13.0)) QJ = QJ*G3

C

```
IF (NX.GE.3 .AND. IZ.GE.59 .AND. NT.GE.50 .AND. N.LT.82)
     ED = E - 3.0
&
B = A
IF (B.LT.35.0) B = 35.0
FX = (12.0 + 0.1*X)*X - (1.0 - 1.0/X**0.5)*B**(2.0/3.0)
ALOGY = (1.0 + 1.5*X)*ALOG(ABS(ED/FX))
IF (ALOGY.GT.10.0) ALOGY=10.0
F1 = 1.0 - EXPF(-EXP(ALOGY))
C = 1.0 + 0.03*X*(A-208.0)
IF (C.LE.1.0) C = 1.0
D = 27.5 - 0.1*(A + (200.0 - A)/X**0.5)
IF ((1.0 - X) * (208.0 - A).GT.0.0) D =
                                   D - 0.03*(1.0 - X)*(208.0 - A)
F = E
IF (NX.GE.3 .AND. IZ.GE.39 .AND. NT.GE.28 .AND. N.LT.50)
      F = E - 3.0
IF (NX.GE.3 .AND. IZ.GE.59 .AND. NT.GE.50 .AND. N.LT.82)
      F = E - 3.0
IF (NX.EQ.1 .AND. IZ.GE.79 .AND. IZ.LE.83) F = E + 5.0
F2 = 3500/C * EXPF(-0.6*X - 0.5*(((F - D) - 5.0*X**1.34)/
       (6.0 - 2.5/X**4))**2)
      * (1.0 - EXPF(-(0.03*A/(2.0 - 1.0/X**4))**3))
G = 0.01*(A - 208.0)
IF (A.LT.208.0) G = 0.0
F3 = (1300.0/(E + 20.0*X**1.5/E))**(1.3 - G)
FH = (1300.0/(EI + 20.0*X**1.5/EI))**(1.3 - G)
FE = F1*(F2 + F3)
IF (E.GE.EI) QJ = FH*QI
IF (E.LT.EI) QJ = FE*QI
IF (Z.EQ.6.0 .AND. (A-X+1.0).EQ.13.0) QJ = QJ*0.4
IF (Z.EQ.3.0 .AND. (A-X+1.0).EQ.9.0) QJ = QJ*0.65
RETURN
END
FUNCTION EXPF(X)
IF (X.LT.-23.0) X = -23.0
IF (X.GT.23.0) X = 23.0
EXPF = EXP(X)
END
```

```
SUBROUTINE ZTABLE (ELOWER, EUPPER, M, IZLO, IZUP, TARGET, PSTEP)
C SUBROUTINE ZTABLE in Module UPROP.FOR
C
C Creates the auxiliary ionization loss data file (ZTABLE.DAT).
C Modified by AJT 5-8-96 to remove examination of old files.
C Parameters
C
         Maximum number of logarithically-spaced energy bins in spectrum
C MARR
C ELOWER Lower energy bound of input and output spectra (>= 0.1 MeV)
C EUPPER Upper energy bound of input and output spectra (<= 100000 MeV)
         Number of logarithmically equally-spaced energy bins (<= MARR)
         Least atomic number of elements transported (>= 1)
C IZLO
         Greatest atomic number of elements transported (<= 109)
C IZUP
C TARGET Name of the target shielding material (>= 12 bytes)
         A small pathlength over which 2 nuclear fragmentations are
         unlikely, typically 0.1 g/cm**2.
С
C
C Important variables
C
         Energy at each grid point after shielding (MeV)
CE
         Stopping power at each grid point after shielding (initially)
C S
         Ratio of stopping powers before and after shielding (modified)
C
         Range at each grid point after shielding (initially)
C R
         Range at each grid point prior to shielding (modified)
         Energy at each grid point prior to shielding
C EP
         Stopping power at each grid point prior to shielding
C SP
C RP
         Range at each energy EP
C Subprograms
C SUBROUTINE RANGE (E, M, Z, A, TARGET, R)
    Returns the range R(M) at M energy grid points E(M) for an element with
   charge Z and mass A in target material TARGET
C FUNCTION STPOW(E, Z, A, TARGET)
    Returns the stopping power STPOW at energy E for an element with charge
С
    Z and mass A in target material TARGET
С
C BLOCK DATA D01
   Defines the atomic masses of elements in the range 1 <= Z <= 109 and
    places them in the array AMASS
C
C
C Data File
C ZTABLE.DAT
   Contains ionization loss data for the transport calculation.
    Automatically created by this subroutine when needed.
PARAMETER (MARR=5000)
      REAL*4 E (MARR), S (MARR), R (MARR), EP (MARR), SP (MARR), RP (MARR)
      CHARACTER*12 TARGET, TARGET$
      INTEGER*4 STAT, CREME96_OPEN
```

COMMON/MASS/AMASS(109)

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```
DATA LMAX, MAXVERS/2,10/
       DATA ELOWER$, EUPPER$, M$, IZLO$, IZUP$, TARGET$,
              PSTEP$/0.,0.,0,0,0,' ',0./
С
       FORMAT Statements
100
       FORMAT(1X,2(1PE10.4,2X),3(I5,2X),A12,2X,1PE10.4)
200
       FORMAT ((1X, 6 (1PE10.4, 2X)))
       OPEN (UNIT=11, FILE='USER: ZTABLE.DAT', STATUS='NEW')
C
       stat = creme96 open('ztable.dat','user',11,'new')
С
       Write header
       ELOWERS=ELOWER
       EUPPER$=EUPPER
      M$=M
       IZLO$=IZLO
       IZUP$=IZUP
       TARGET$=TARGET
       PSTEP$=PSTEP
       WRITE (11,100) ELOWER, EUPPER, M, IZLO, IZUP, TARGET, PSTEP
       WRITE (11, '(A)') ''
C
      Compute vector of energies
      DE= (EUPPER/ELOWER) ** (1./FLOAT (M-1))
      E(1)=ELOWER
      DO J=2, M-1
         E(J) = E(J-1) *DE
      END DO
      E(M)=EUPPER
C
      Compute parameters
      DO J=IZLO, IZUP
         Z=FLOAT(J)
         A=AMASS(J)
         CALL RANGE(E,M,Z,A,TARGET,R)
         DO K=1, M
           S(K) = STPOW(E(K), Z, A, TARGET)
         END DO
         DO K=1.M
           DO KK=K, M
             IF (R(KK).GE.R(K)+PSTEP) GOTO 300
           END DO
300
           EP(K) = E(KK) - (R(KK) - R(K) - PSTEP) *S(KK)
           R(K) = R(K) + PSTEP
         END DO
C
         Iterate LMAX times to improve estimate of EP
         DO L=1, LMAX
           CALL RANGE (EP, M, Z, A, TARGET, RP)
           DO K=1,M
             SP(K) = STPOW(EP(K), Z, A, TARGET)
             EP(K) = EP(K) - (RP(K) - R(K)) *SP(K)
           END DO
         END DO
```

```
Compute ratio of stopping powers
C
        DO K=1, M
          S(K) = SP(K)/S(K)
        END DO
C
        Write output to ZTABLE.DAT
        WRITE (11,200) (EP(K), K=1, M)
        WRITE (11,100)
        WRITE (11,200) (S(K),K=1,M)
        WRITE (11,100)
      END DO
      Close output file and stop
С
      CLOSE (UNIT=11)
      RETURN
      END
```

```
CHARACTER*12 NAME
                               REAL NA(28), IADJ(28), NASPM(28), DENS, ETAD
                               INTEGER NZ (28), IGAS, NAS, STAT, CREME96_OPEN
                               COMMON /TBLOCK/DENS, ETAD, IGAS, NAS,
                                                                          NZ, NA, IADJ, NASPM,
                       &
                                                                          NTOTAL, AVGZ, AVGZ2, AVGA, AVGI
                       &
                               Check to see if current target is LNAME
         !
                               IF (LNAME.EQ.NAME) RETURN
                               Open TARGET.DAT and read data for LNAME
                               OPEN (UNIT=10, FILE='CREME96: TARGET.DAT',
         С
                          & STATUS='OLD', READONLY, SHARED)
         С
                          stat = creme96_open('target.dat','cr96tables',10,'old')
                               FORMAT(1X, I3)
           1
                               FORMAT (1X, A12, 2X, F9.6, 2X, F9.6, 2X, I1, 2X, I2)
           2
                               FORMAT (1X, I3, 2X, F8.4, 2X, F5.1, 2X, F9.5)
           3
                               READ(10,1) NM
                               DO J1=1, NM
                                     READ(10,2) NAME, DENS, ETAD, IGAS, NAS
                                     DO J2=1, NAS
                                           READ(10,3) NZ(J2), NA(J2), IADJ(J2), NASPM(J2)
The state of the s
                                     END DO
                                     IF (LNAME.EQ.NAME) THEN
                                           CLOSE (UNIT=10)
                                           GO TO 100
L
                                     ENDIF
1
                                END DO
32
                                CLOSE (UNIT=10)
فعظ
                                PRINT *,' ***** Target Data not available *****'
1
                               STOP
100
                               CONTINUE
Ē.
Ţ
                                Compute material parameters
                               NTOTAL=0
                                AVGZ=0.
                                AVGZ2=0.
                                AVGA=0.
                                AVGI=0.
                               DO J1=1, NAS
                                     NTOTAL=NTOTAL + NASPM(J1)
                                      AVGZ=AVGZ + NASPM(J1)*FLOAT(NZ(J1))
                                      AVGZ2=AVGZ2 + NASPM(J1)*FLOAT(NZ(J1))**2
                                      AVGA=AVGA + NASPM(J1)*NA(J1)
                                      AVGI=AVGI + NASPM(J1)*ALOG(IADJ(J1))
                                END DO
                                AVGZ=AVGZ/FLOAT (NTOTAL)
                                AVGZ2=AVGZ2/FLOAT(NTOTAL)
                                AVGA=AVGA/FLOAT (NTOTAL)
                                AVGI=EXP(AVGI/FLOAT(NTOTAL))
                                RAT=AVGZ/AVGA
```

RETURN END

SUBROUTINE ZTARGET (LNAME)

INCLUDE 'CREME96: ZCOMMON.CMN'

CHARACTER*12 LNAME

C

CREME96 ROADMAP For orbit inside Earth's For orbits at or beyond Auxiliary Programs you may magnetosphere, start here: geosynchronous, start here: wish to run: GTRN: 200 Evaluates geomagnetic shielding 100 FLUX: Evaluates fluxes at external surface of spacecraft TRANS: SHIELDFILE: Formats user-supplied Transports nuclei through shielding distribution for shielding input to TRANS LET: 40 PUP: Calculates Proton-Induced Linear Energy SEU Rates Transfer (LET) Spectra HUP: 700 Finds Finds Direct-Ionization Induced SEU L Rates W.

F16. I Figur 10.

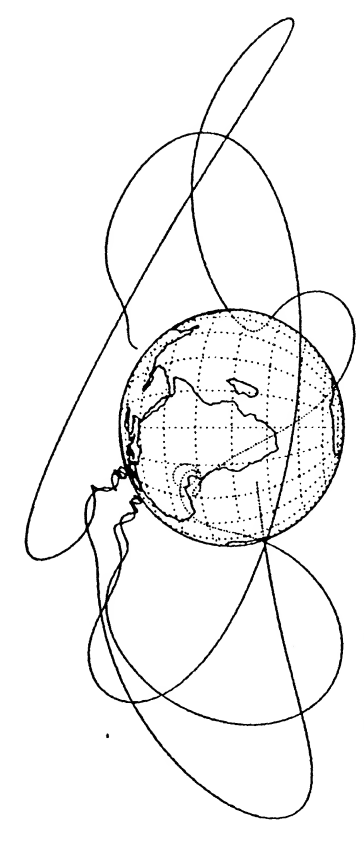


Fig. 5.1 - This figure shows the computed trajectory of a cosmic ray in the earth's magnetic field. Such complex trajectories are not unusual for cosmic rays in the penumbral shadow near the Stormer cutoff (Lund, 1980).

F16.2

Choose number of steps per each orbit (N_{spo}) of the Earth (e.g. $N_{spo} = 200$ steps), set orbit step counter N = 0 and number transmitted $N_{tr} = 0$, and specify required accuracy of numerical integration (typically 10⁻⁵).

310

Input orbital parameters: inclination, apogee, perigee, initial time (to), orbital duration (T), initial displacement of perigee from ascending mode, initial longitude of ascending node, and initial displacement from the ascending mode.

Calculate orbital period (P)

N

W.

from input orbital parameters using standard orbit generation routines (e.g. Adams et al., 1986). Calculate:

 $N_{max} = N_{spo} M_{adpos} T/P$ and $\delta t = P/N_{spo}$.

320

Find spacecraft's orbital location for time $t = t_0 + N \delta t$ using numerical solution of standard equations for orbit generator, e.g. those given on pp. 43 - 48 of (Sterne, 1960), as implemented in the CREME orbit generator (Adams et al., 1986). Set particle trajectory time t' = t and time step $\delta t' = 0$. Set M = 0.

Differential equation for integrating is the Lorenz equation in magnetic field, using "backward" solution (O replaced by -Q, t' replaced by -t') of:

$$F = m \gamma dv/dt' = Q v x B$$

$$\begin{split} \mathbf{B} &= \mathbf{B}_{\text{IGRF}}(\mathbf{r},\,t') \, + \mathbf{B}_{\text{Tsyg89}}(\text{Kp,r,}\,t') \\ &+ \, \delta \, \mathbf{B}_{\text{extended Tsyg89}}(\text{Kp,Dst,}\,\mathbf{r,}t') \end{split}$$

NOTE: In the "backward" solution, the actual particle's final position is the initial position along the numerical integration of the "backward" particle trajectory, and the "backward" particle's initial velocity is in the opposite direction as the actual particle's final velocity.

See text for description of B-field calculations.

Set r = particle's final position and v= particle's final velocity vector. These are determined from the spacecraft's latitude, longitude, and altitude, combined with the particle's rigidity, final θ , and final φ .

Generate particle's final heta and

 φ for each orbital location, using random number techniques, i.e. x1 and x2 are two distinct outputs from a random number generator.

$$\cos \theta = \cos \theta_{\text{max}} + x_1 (\cos \theta_{\text{min}} - \cos \theta_{\text{max}})$$

$$\varphi = \varphi_{\min} + x_2 (\varphi_{\max} - \varphi_{\min})$$

M = M + 1

NOTE: many modern computers have built in random number generator functions. For computers which do not, one can use the subroutines for random number generation from (Press et al., 1992).

Evaluate B-field and d v/dt at location r and time t' along the particle's trajectory, i.e. t' is the time along the particle's trajectory.

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Specifically, perform Bulirsch-Stoer numerical integration, using formulae & algorithms as presented on pp. 718-725 of (Press et al., 1992). The outputs are the particle's radial position r, velocity v, and time $t' = t + \delta t'$ after the numerical integration step; and an estimate of $\delta t'$ for the next particle-trajectory integration time-step.

NOTE: δ t' is re-calculated within each execution of the adaptive Bulirsch-Stoer integration.

Evaluate if particle trajectory encountered solid Earth (forbidden), entered atmosphere (forbidden), crossed magnetospheric boundary (transmitted), or still within magnetosphere.

If transmitted, $N_{tr} = N_{tr} + 1$

If forbidden, $N_{fb} = N_{fb} + 1$

If within magnetosphere.

If transmitted or forbidden,

N = N + 1

If $M < M_{adpos}$ 370 If $N \leq N_{max}$

If $N > N_{max}$

Calculate Geomagnetic Transmission (GT) for this rigidity (R) and output the result:

 $GT(R) = N_{tr} / N_{max}$

NOTE: $N_{fb} = N_{max} - N_{tr}$

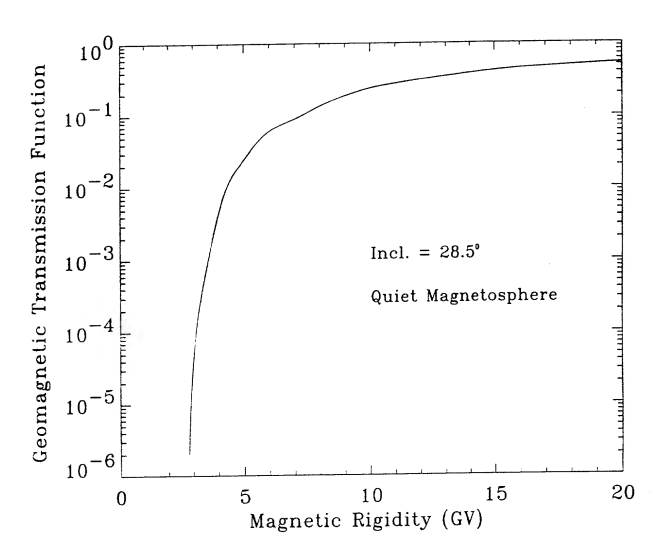
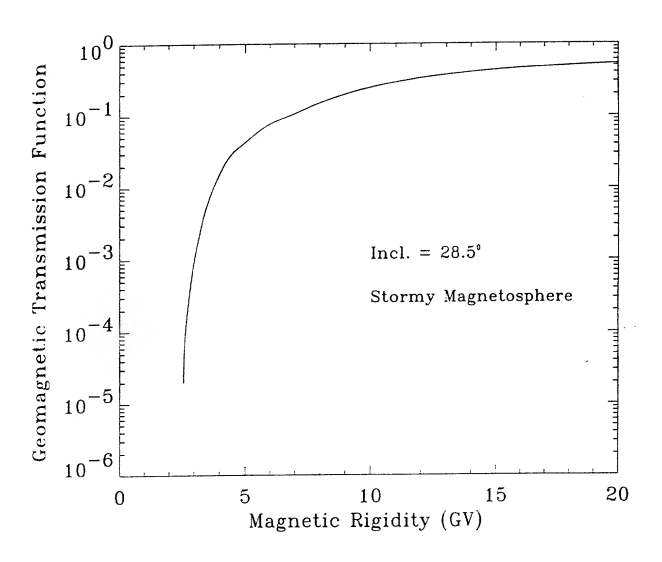
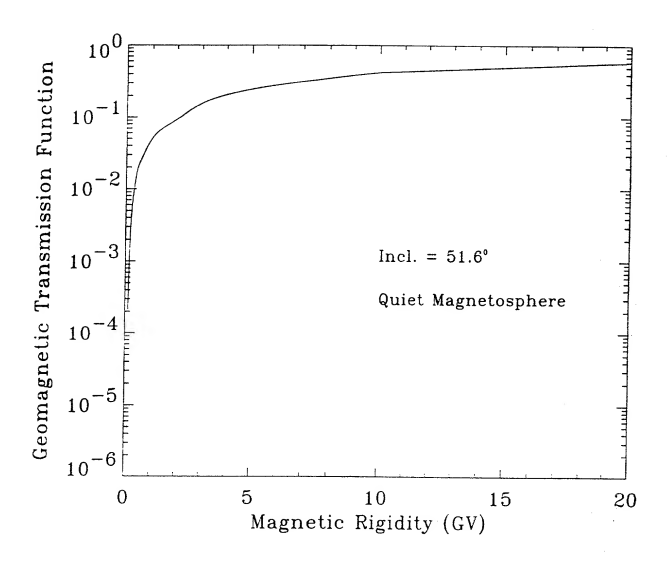


Figure 1



F16.5



F16. 6

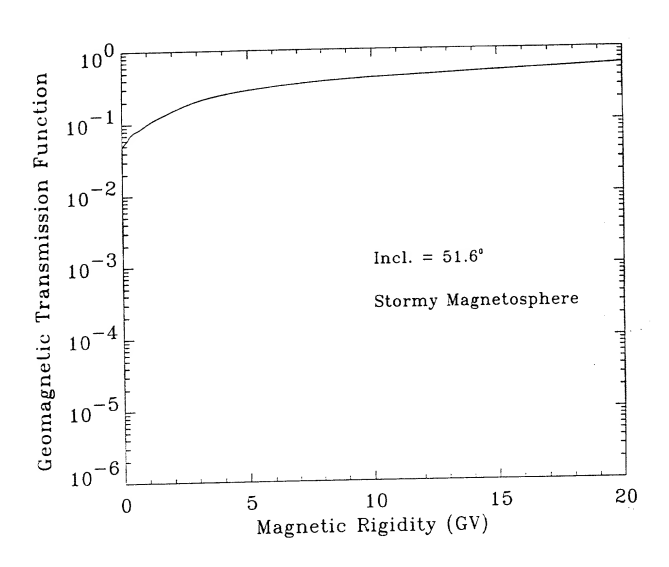
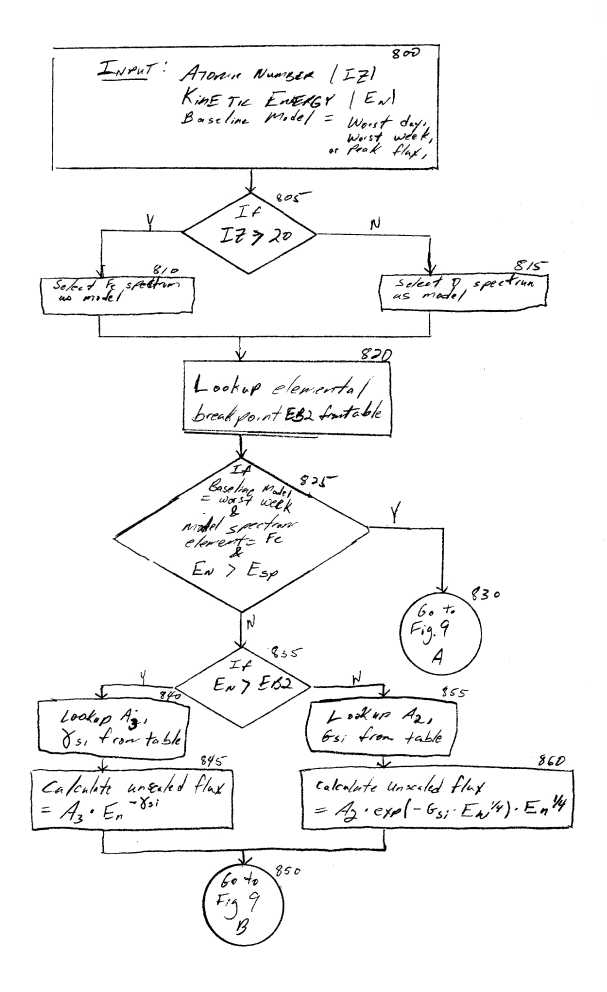
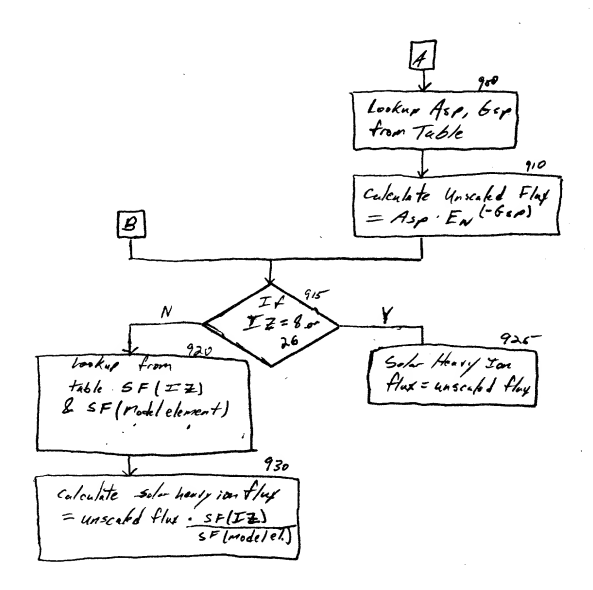


FIG. 7





CREME96 Main Menu

See brief instructions at the bottom of this page. See How to Run CREME96 for more details.

GO	RESET FORM

Routine	<u>User Request File</u> list	Edit User Request File	Run Routine
TPS	■ .	0	
GTRN	T	0	
FLUX	7	0	
TRANS	7	0	
LETSPEC	V	0	
PUP		0	G
HUP	V	0	
UTILTIES	Г		

- 1. To CREATE a User Request File:
 - o Click on the Edit button by the corresponding routine, leaving the User Request File blank. (If necessary, use the blank space in the pulldown menu to clear a name already typed there.)
 - o Click on the "GO" bar.
- 2. To EDIT an existing User Request File:
 - o Click on the Edit button by the corresponding routine.
 - o Select a User Request File from the appropriate pull-down menu.
 - o Click on the "GO" bar.
- 3. To RUN one or more routines:
 - o If an Edit button has been clicked "on", use the RESET bar to clear it.
 - o Select User Request File(s) from the appropriate pull-down menu(s).
 - o Click on the RUN button(s) in that line(s)
 - o Click on the "GO" bar.
- 4. To access CREME96 Utilities:
 - o Click on the RUN button on that line
 - o Click on the "GO" bar.

Note: You can create or edit only one User Request File at a time. But you can submit several files for running (sequentially, in the order shown in the table) by ehecking more than one "RUN" button and then clicking on the "GO" bar.

Questions? Comments? Send mail to tylka@crs2.nrl.navy.mil.

You may terminate your CREME96 session at any time.

F16.10

Lyw 3

AND POWER OF ATTORNEY

As a below named inventor, I hereby declare that: My residence, post office address and citizenship are as stated below next to my name. I believe I am the original, first, and joint inventor of the subject matter which is claimed and for which a patent is sought on the invention entitled: METHOD AND APPARATUS FOR MODELING COSMIC RAY EFFECTS ON MICROELECTRONICS, the specification of which is attached hereto. I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above. I acknowledge the duty to disclose information which is material to the examination of this application in accordance with Title 37, Code of Federal Regulations, \$1.56(a).

I hereby claim foreign priority benefits under Title 35, United States Code, \$119 of any foreign

I hereby claim foreign prio applications for patent or application for patent or i which priority is claimed:	inventor's ce	rtificate lis	ted below and have	also ideni	tified below any foreign
Number	Country		Filing Date		Priority (Yes/No)
I hereby claim the benefit below and, insofar as the s prior United States applica Code, \$112, I acknowledge t Regulations, \$1.56(a) which PCT international filing da	ubject matter tion in the r he duty to di occurred bet	r of each of t manner provide isclose materi tween the fili	he claims of this a d by the first para al information as d	pplication graph of efined in	title 35, United States Title 37, Code of Federal
U.S. Appl. Serial No.	U.	S. Filing Dat	е	Status (patente	ed/pending/abandoned)
POWER OF ATTORNEY: As a na prosecute this application and hereby certify that the application: Thomas E. McDonnell, Reg. N	and transact Government	all business of the United	in the Patent and T States has the irre	rademark vocable r	Office connected therewith,
SEND CORRESPONDENCE TO: Associate Counsel (Paten Naval Research Laborator Washington, D.C. 20375-50	У	8.2	DIRECT TELEPHONE John J. Karasek Reg. No. 36,182 (202)404-1552		
I hereby declare that all son information and belief a knowledge that willful falsooth, under Section 1001 of peopardize the validity of	re believed se statements Title 18 of	to be true; an and the like the United St	d further that thes so made are punisha ates Code, and that	e stateme ble by fi such wil	nts were made with the ne or imprisonment, or
Full name of joint inventor	: 1: James H	. Adams			
Inventor's signature:				DATE:	
Residence: Citizenship: Post Office Address:					
Full name of joint inventor	2: Paul B	oberg	•		11-21-97
Full name of joint inventor Inventor's signature:	aul.	K. ZEK	<u> </u>	DATE:	12111
Residence: College Park, N Citizenship: USA Post Office Address: 5023	10 .		•		
Full name of joint inventor	c 3: Buddy	Brownstein	4		- 1- 10-
Inventor's signature:	uddy	Pround		DATE:	12/31/97
Residence: Oxen Hill, MD Citizenship: USA Post Office Address: 7402	0				
Full name of joint inventor	r 4: Willia	m Dietrich			
Inventor's signature:				DATE:	

Inventor's signature: ___

Citizenship: Post Office Address:

FILING DECLARATION AND POWER OF ATTORNEY

Navy Case No. 78,824 14/3/97

Full name of joint inventor 5: Erwin Flueckiger		
Inventor's signature:	DATE:	
Residence: Citizenship: Post Office Address:		
Full name of joint inventor 6: Edward Petersen		
Inventor's signature:	DATE:	
Residence: Citizenship: Post Office Address:		
Full name of joint inventor 7: Margaret Shea		
Inventor's signature:	DATE:	
Residence: Citizenship: Post Office Address:		
Full name of joint inventor 8: Don Smart		
Inventor's signature:	DATE:	
Residence: Citizenship: Post Office Address:		
Full name of joint inventor 9: Edward Smith		
Inventor's signature:	_ DATE:	
Residence: Citizenship: Post Office Address:		
@Full name of joint inventor 10: Allan Tylka		
Inventor's signature:	_ DATE:	
Residence:		